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# Example Programs for KINSOL v2.2.0

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# Example Programs for KINSOL v2.2.0

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# 1 Introduction

This report is intended to serve as a companion document to the User Documentation of KINSOL [1]. It provides details, with listings, on the example programs supplied with the KINSOL distribution package.

The KINSOL distribution contains examples of four types: serial C examples, parallel C examples, and serial and parallel FORTRAN examples. The following lists summarize all of these examples. Supplied in the `sundials/kinsol/examples_ser` directory is the following serial example (using the `NVECTOR_SERIAL` module):

- `kinwebs` is a demonstration program for KINSOL with the Krylov linear solver.

This program solves a nonlinear system that arises from a system of partial differential equations. The PDE system is a six-species food web population model, with predator-prey interaction and diffusion on the unit square in two dimensions.

The preconditioner matrix is a block-diagonal matrix based on the partial derivatives of the interaction terms only.

Supplied in the `sundials/kinsol/examples_par` directory are the following two parallel examples (using the `NVECTOR_PARALLEL` module):

- `kinwebp` is a parallel implementation of `kinwebs`.
- `kinwebbpd` solves the same problem as `kinwebp`, with a block-diagonal matrix with banded blocks as a preconditioner, generated by difference quotients, using the module `KINBBDPRE`.

With the FKINSOL module, in the directories `sundials/kinsol/fcmix/examples_ser` and `sundials/kinsol/fcmix/examples_par`, are the following examples for the FORTRAN-C interface:

- `kindiagsf` solves a nonlinear system of the form  $u_i^2 = i^2$  using an approximate diagonal preconditioner.
- `kindiagpf` is a parallel implementation of `kindiagsf`.

In the following sections, we give detailed descriptions of some (but not all) of these examples. The Appendices contain complete listings of those examples described below. We also give our output files for each of these examples, but users should be cautioned that their results may differ slightly from these. Differences in solution values may differ within the tolerances, and differences in cumulative counters, such as numbers of steps or Newton iterations, may differ from one machine environment to another by as much as 10% to 20%.

In the descriptions below, we make frequent references to the KINSOL User Document [1]. All citations to specific sections (e.g. §5.1) are references to parts of that User Document, unless explicitly stated otherwise.

**Note.** The examples in the KINSOL distribution are written in such a way as to compile and run for any combination of configuration options during the installation of SUNDIALS (see §2). As a consequence, they contain portions of code that will not be typically

present in a user program. For example, all C example programs make use of the variable `SUNDIALS_EXTENDED_PRECISION` to test if the solver libraries were built in extended precision and use the appropriate conversion specifiers in `printf` functions. Similarly, the FORTRAN examples in FKINSOL are automatically pre-processed to generate source code that corresponds to the precision in which the KINSOL libraries were built (see §3 in this document for more details).

## 2 C example problems

### 2.1 A serial example: `kinwebs`

We give here an example that illustrates the use of KINSOL with the Krylov method SPGMR, in the `KINSPGMR` module, as the linear system solver. The source file, `kinwebs.c`, is listed in Appendix A.

This program solves a nonlinear system that arises from a discretized system of partial differential equations. The PDE system is a six-species food web population model, with predator-prey interaction and diffusion on the unit square in two dimensions. Given the dependent variable vector of species concentrations  $c = [c_1, c_2, \dots, c_{n_s}]^T$ , where  $n_s = 2n_p$  is the number of species and  $n_p$  is the number of predators and of prey, then the PDEs can be written as

$$d(i) \cdot \left( \frac{\partial^2 c_i}{\partial x^2} + \frac{\partial^2 c_i}{\partial y^2} \right) + f_i(x, y, c) = 0 \quad (i = 1, \dots, n_s), \quad (1)$$

where the subscripts  $i$  are used to distinguish the species, and where

$$f_i(x, y, c) = c_i \cdot \left( b(i) + \sum_{j=1}^{n_s} a(i, j) \cdot c_j \right). \quad (2)$$

The problem coefficients are given by

$$a_{ij} = \begin{cases} -1 & i = j \\ -0.5 \cdot 10^{-6} & i \leq n_p, j > n_p \\ 10^4 & i > n_p, j \leq n_p \\ 0 & \text{all other ,} \end{cases}$$

$$b_i = b_i(x, y) = \begin{cases} 1 + \alpha xy & i \leq n_p \\ -1 - \alpha xy & i > n_p, \end{cases}$$

and

$$d_i = \begin{cases} 1 & i \leq n_p \\ 0.5 & i > n_p. \end{cases}$$

The spatial domain is the unit square  $(x, y) \in [0, 1] \times [0, 1]$ .

Homogeneous Neumann boundary conditions are imposed and the initial guess is constant in both  $x$  and  $y$ . For this example, the equations (1) are discretized spatially with standard central finite differences on a  $8 \times 8$  mesh with  $n_s = 6$ , giving a system of size 384.

Among the initial `#include` lines in this case are lines to include `kinspgmr.h` and `sundialsmath.h`. The first contains constants and function prototypes associated with the SPGMR method. The inclusion of `sundialsmath.h` is done to access the `MAX` and `ABS` macros, and the `RSqrt` function to compute the square root of a `realtype` number.

The `main` program calls `KINCreate` and then calls `KINMalloc` with the name of the user-supplied system function `func` and solution vector as arguments. The `main` program then calls a number of `KINSet*` routines to notify KINSOL of the function data pointer, the positivity constraints on the solution, and convergence tolerances on the system function and step size. It calls `KINSpgrm` (see §5.4.2) to specify the `KINSPGMR` linear solver, and passes a value of 15 as the maximum Krylov subspace dimension, `maxl`. Next, a

maximum value of `maxlrst = 2` restarts is imposed and the user-supplied preconditioner setup and solve functions, `PrecSetupBD` and `PrecSolveBD`, are specified through calls to `KINSpgmrSetPrecSetupFn` and `KINSpgmrSetPrecSolveFn`, respectively (see §5.4.4). The data pointer passed to `KINSpgmrSetPrecData` is passed to `PrecSetupBD` and `PrecSolveBD` whenever these are called.

Next, `KINSol` is called, the return value is tested for error conditions, and the approximate solution vector is printed via a call to `PrintOutput`. After that, `PrintFinalStats` is called to get and print final statistics, and memory is freed by calls to `N_VDestroy_Serial`, `FreeUserData` and `KINFree`. The statistics printed are the total numbers of nonlinear iterations (`nni`), of `func` evaluations (excluding those for  $Jv$  product evaluations) (`nfe`), of `func` evaluations for  $Jv$  evaluations (`nfeSG`), of linear (Krylov) iterations (`nli`), of preconditioner evaluations (`npe`), and of preconditioner solves (`nps`). All of these optional outputs and others are described in §5.4.5.

Mathematically, the dependent variable has three dimensions: species number,  $x$  mesh point, and  $y$  mesh point. But in `NVECTOR_SERIAL`, a vector of type `N_Vector` works with a one-dimensional contiguous array of data components. The macro `IJ_Vptr` isolates the translation from three dimensions to one. Its use results in clearer code and makes it easy to change the underlying layout of the three-dimensional data. Here the problem size is 384, so we use the `NV_DATA_S` macro for efficient `N_Vector` access. The `NV_DATA_S` macro gives a pointer to the first component of a serial `N_Vector` which is then passed to the `IJ_Vptr` macro.

The preconditioner used here is the block-diagonal part of the true Newton matrix and is based only on the partial derivatives of the interaction terms  $f$  in (2) and hence its diagonal blocks are  $n_s \times n_s$  matrices ( $n_s = 6$ ). It is generated and factored in the `PrecSetupBD` routine and backsolved in the `PrecSolveBD` routine. See §5.5.4 for detailed descriptions of these preconditioner functions.

The program `kinwebs.c` uses the “small” dense functions for all operations on the  $6 \times 6$  preconditioner blocks. Thus it includes `smalldense.h`, and calls the small dense matrix functions `denalloc`, `denallocpiv`, `denfree`, `denfreepiv`, `gefa`, and `gesl`. The small dense functions are generally available for KINSOL user programs (for more information, consult the comments in the header file `smalldense.h`).

In addition to the functions called by KINSOL, `kinwebs.c` includes definitions of several private functions. These are: `AllocUserData` to allocate space for  $P$  and the pivot arrays; `InitUserData` to load problem constants in the `data` block; `FreeUserData` to free that block; `SetInitialProfiles` to load the initial values in `cc`; `PrintOutput` to retrieve and print selected solution values; `PrintFinalStats` to print statistics; and `check_flag` to check return values for error conditions.

The output generated by `kinwebs` is shown below.

```
----- kinwebs sample output -----  
  
Predator-prey test problem -- KINSol (serial version)  
  
Mesh dimensions = 8 X 8  
Number of species = 6  
Total system size = 384  
  
Flag globalstrategy = 1 (1 = Inex. Newton, 2 = Linesearch)  
Linear solver is SPGMR with maxl = 15, maxrst = 2
```

```

Preconditioning uses interaction-only block-diagonal matrix
Positivity constraints imposed on all components
Tolerance parameters: fnormtol = 1e-07 scsteptol = 1e-13

Initial profile of concentration
At all mesh points: 1 1 1 30000 30000 30000

Computed equilibrium species concentrations:

At bottom left:
1.16428 1.16428 1.16428 34927.5 34927.5 34927.5

At top right:
1.25797 1.25797 1.25797 37736.7 37736.7 37736.7

Final Statistics..

nni      =      7      nli      =    230
nfe      =      8      nfeSG   =    237
nps      =    237      npe      =      1      ncfl   =      4

```

## 2.2 A parallel example: `kinwebbd`

In this example, `kinwebbd`, we solve the same problem as with `kinwebs` above, but in parallel, and instead of supplying the preconditioner we use the KINBBDPRE module, which generates and uses a band-block-diagonal preconditioner. The upper and lower half-bandwidths of the Jacobian block on each process are both equal to  $2 \cdot \text{NUM\_SPECIES} - 1$ , and that is the value supplied as `mu` and `ml` in the call to `KINBBDPrecAlloc`.

In this case, we think of the parallel MPI processes as being laid out in a rectangle, and each process being assigned a subgrid of size `MXSUB` $\times$ `MYSUB` of the  $x - y$  grid. If there are `NPEX` processes in the  $x$  direction and `NPEY` processes in the  $y$  direction, then the overall grid size is  $\text{MX} \times \text{MY}$  with  $\text{MX} = \text{NPEX} \times \text{MXSUB}$  and  $\text{MY} = \text{NPEY} \times \text{MYSUB}$ , and the size of the ODE system is  $\text{NUM\_SPECIES} \cdot \text{MX} \cdot \text{MY}$ .

The evaluation of the nonlinear system function is performed in `func`. In this parallel setting, the processes first communicate the subgrid boundary data and then compute the local components of the nonlinear system function. The MPI communication is isolated in the private function `ccomm` (which in turn calls `BRecvPost`, `BSend`, and `BRecvWait`) and the subgrid boundary data received from neighboring processes is loaded into the work array `cext`. The computation of the nonlinear system function is done in `func_local` which starts by copying the local segment of the `cc` vector into `cext` and then by imposing the boundary conditions by copying the first interior mesh line from `cc` into `cext`. After this, the nonlinear system function is evaluated by using central finite-difference approximations using the data in `cext` exclusively.

The function `func_local` is also passed as the `gloc` argument to `KINBBDPrecAlloc`. Since all communication needed for the evaluation of the local approximation of  $f$  used in building the band-block-diagonal preconditioner is already done for the evaluation of  $f$  in `func`, a NULL pointer is passed as the `gcomm` argument to `KINBBDPrecAlloc`.

The **main** program resembles closely that of the **kinwebs** example, with particularization arising from the use of the parallel MPI NVECTOR\_PARALLEL module. It begins by initializing MPI and obtaining the total number of processes and the id of the local process. The local length of the solution vector is then computed as **NUM\_SPECIES****·MXSUB****·MYSUB**. Distributed vectors are created by calling the constructor defined in NVECTOR\_PARALLEL with the MPI communicator and the local and global problem sizes as arguments. All output is performed only from the process with id equal to 0. Finally, after all memory deallocation, the MPI environment is terminated by calling **MPI\_Finalize**.

The output generated by **kinwebbbd** is shown below.

```
----- kinwebbbd sample output -----
Predator-prey test problem-- KINSol (parallel-BBD version)

Mesh dimensions = 20 X 20
Number of species = 6
Total system size = 2400

Subgrid dimensions = 10 X 10
Processor array is 2 X 2

Flag globalstrategy = 1 (1 = Inex. Newton, 2 = Linesearch)
Linear solver is SPGMR with maxl = 20, maxlrst = 2
Preconditioning uses band-block-diagonal matrix from KINBBDPRE
with matrix half-bandwidths ml, mu = 11 11
Tolerance parameters: fnormtol = 1e-07 scsteptol = 1e-13

Initial profile of concentration
At all mesh points: 1 1 1 30000 30000 30000

Computed equilibrium species concentrations:

At bottom left:
1.165 1.165 1.165 34949 34949 34949

At top right:
1.25552 1.25552 1.25552 37663.2 37663.2 37663.2

Final Statistics..

nni      =     10    nli      =    540
nfe      =     11    nfeSG =    550
nps      =    550    npe      =     1    ncfl      =     7
```

### 3 Fortran example problems

The FORTRAN example problem programs supplied with the KINSOL package are all written in standard F77 Fortran and use double-precision arithmetic. However, when the FORTRAN examples are built, the source code is automatically modified according to the configure options supplied by the user and the system type. Integer variables are declared as `INTEGER*n`, where  $n$  denotes the number of bytes in the corresponding C type (`long int` or `int`). Floating-point variable declarations remain unchanged if double-precision is used, but are changed to `REAL*n`, where  $n$  denotes the number of bytes in the SUNDIALS type `realtype`, if using single-precision. Also, if using single-precision, declarations of floating-point constants are appropriately modified, e.g.; `0.5D-4` is changed to `0.5E-4`.

The two examples supplied with the `FKINSOL` module are very simple tests of the FORTRAN-C interface module. They solve the nonlinear system

$$F(u) = 0, \quad \text{where } f_i(u) = u_i^2 - i^2, 1 \leq i \leq N.$$

#### 3.1 A serial example: `kindiagsf`

The `kindiagsf` program (for which the source code is listed in Appendix C) solves the above problem using the serial `NVECTOR_SERIAL` module.

The main program begins by calling `fnavinits` to initialize computations with the serial `NVECTOR_SERIAL` module. Next, the array `uu` is set to contain the initial guess  $u_i = 2i$ , the array `scale` is set with all components equal to 1.0 (meaning that no scaling is done), and the array `constr` is set with all components equal to 0.0 to indicate that no inequality constraints should be imposed on the solution vector.

The KINSOL solver is initialized and memory for it is allocated by calling `fkinmalloc`, which also specifies the maximum number of iterations between calls to the preconditioner setup routine (`msbpre = 5`), the tolerance for stopping based on the function norm (`fnormtol = 10^{-5}`), the tolerance for stopping based on the step length (`scsteptol = 10^{-4}`), and that no optional inputs are provided (`inopt = 0`).

Next, the KINSPGMR linear solver module is attached to KINSOL by calling `fkinspgmr`, which also specifies the maximum Krylov subspace dimension (`maxl = 10`) and the maximum number of restarts allowed for SPGMR (`maxrst = 2`). The KINSPGMR module is directed to use the supplied preconditioner by calling the routines `fkinspgmrsetpsol` and `fkinspgmrsetpset` with a first argument equal to 1. The solution of the nonlinear system is obtained after a successful return from `fkinsol`, which is then printed to unit 6 (stdout).

Memory allocated for the KINSOL solver is released by calling `fkinfree` and computations with the `NVECTOR_SERIAL` module are terminated by calling `fnavfrees`.

The user-supplied routine `fckfun` contains a straightforward transcription of the nonlinear system function  $f$ , while the routine `fkpset` sets the array `pp` (in the common block `pcom`) to contain an approximation to the reciprocals of the Jacobian diagonal elements. The components of `pp` are then used in `fkpset` to solve the preconditioner linear system  $Px = v$  through simple multiplications.

The following is sample output from `kindiagsf`, using  $N = 128$ .

```
kindiagsf sample output
Example program kindiagsf:
This fkinsol example code solves a 128 eqn diagonal algebraic system.
```

Its purpose is to demonstrate the use of the Fortran interface in a serial environment.

```
globalstrategy = KIN_INEXACT_NEWTON
```

```
FKINSOL return code is 0
```

The resultant values of uu are:

1	1.000000	2.000000	3.000000	4.000000
5	5.000000	6.000000	7.000000	8.000000
9	9.000000	10.000000	11.000000	12.000000
13	13.000000	14.000000	15.000000	16.000000
17	17.000000	18.000000	19.000000	20.000000
21	21.000000	22.000000	23.000000	24.000000
25	25.000000	26.000000	27.000000	28.000000
29	29.000000	30.000000	31.000000	32.000000
33	33.000000	34.000000	35.000000	36.000000
37	37.000000	38.000000	39.000000	40.000000
41	41.000000	42.000000	43.000000	44.000000
45	45.000000	46.000000	47.000000	48.000000
49	49.000000	50.000000	51.000000	52.000000
53	53.000000	54.000000	55.000000	56.000000
57	57.000000	58.000000	59.000000	60.000000
61	61.000000	62.000000	63.000000	64.000000
65	65.000000	66.000000	67.000000	68.000000
69	69.000000	70.000000	71.000000	72.000000
73	73.000000	74.000000	75.000000	76.000000
77	77.000000	78.000000	79.000000	80.000000
81	81.000000	82.000000	83.000000	84.000000
85	85.000000	86.000000	87.000000	88.000000
89	89.000000	90.000000	91.000000	92.000000
93	93.000000	94.000000	95.000000	96.000000
97	97.000000	98.000000	99.000000	100.000000
101	101.000000	102.000000	103.000000	104.000000
105	105.000000	106.000000	107.000000	108.000000
109	109.000000	110.000000	111.000000	112.000000
113	113.000000	114.000000	115.000000	116.000000
117	117.000000	118.000000	119.000000	120.000000
121	121.000000	122.000000	123.000000	124.000000
125	125.000000	126.000000	127.000000	128.000000

Final statistics:

```
nni =    7, nli =   21, nfe =     8, npe =     2, nps =   28, ncfl =    0
```

### 3.2 A parallel example: kindiagpf

The program `kindiapf` (listed in Appendix D) is a straightforward modification of `kindiagsf` to use the parallel MPI `NVECTOR_PARALLEL` module.

After initialization of MPI, the NVECTOR\_PARALLEL module is initialized by calling `fnpinitp` with the local and global vector sizes as its first two arguments. The problem set-up (KINSOL initialization, KINSPGMR specification) and solution steps are the same as in `kindiagsf`. Upon successful return from `fkinsol`, the solution segment local to the process with id equal to 0 is printed to the screen. Finally, the KINSOL memory is released, NVECTOR\_PARALLEL computations are finalized, and the MPI environment is terminated.

For this simple example, no inter-process communication is required to evaluate the nonlinear system function  $f$  or the preconditioner. As a consequence, the user-supplied routines `fkfun`, `fkpset`, and `fkpssol` are basically identical to those in `kindiagsf`.

Sample output from `kindiapf`, for  $N = 128$ , follows.

```
kindiagpf sample output
Example program kindiagpf:

This fkinsol example code solves a 128 eqn diagonal algebraic system.
Its purpose is to demonstrate the use of the Fortran interface
in a parallel environment.

globalstrategy = KIN_INEXACT_NEWTON
FKINSOL return code is      0

The resultant values of uu (process 0) are:
   1   1.000000   2.000000   3.000000   4.000000
   5   5.000000   6.000000   7.000000   8.000000
   9   9.000000  10.000000  11.000000  12.000000
  13  13.000000  14.000000  15.000000  16.000000
  17  17.000000  18.000000  19.000000  20.000000
  21  21.000000  22.000000  23.000000  24.000000
  25  25.000000  26.000000  27.000000  28.000000
  29  29.000000  30.000000  31.000000  32.000000

Final statistics:
nni =     7,  nli =    21,  nfe =     8,  npe =     2,  nps=  28,  ncfl=    0
```

## References

- [1] A. M. Collier, A. C. Hindmarsh, R. Serban, and C.S. Woodward. User Documentation for KINSOL v2.2.0. Technical Report UCRL-SM-208116, LLNL, 2004.

## A Listing of kinwebs.c

```
1  /*
2  * -----
3  * $Revision: 1.14 $
4  * $Date: 2004/11/15 17:28:14 $
5  * -----
6  * Programmer(s): Allan Taylor, Alan Hindmarsh and
7  *                 Radu Serban @ LLNL
8  * -----
9  * Example (serial):
10 *
11 * This example solves a nonlinear system that arises from a system
12 * of partial differential equations. The PDE system is a food web
13 * population model, with predator-prey interaction and diffusion
14 * on the unit square in two dimensions. The dependent variable
15 * vector is the following:
16 *
17 *      1   2           ns
18 * c = (c , c , ... , c )      (denoted by the variable cc)
19 *
20 * and the PDE's are as follows:
21 *
22 *          i           i
23 *          0 = d(i)*(c + c ) + f (x,y,c)  (i=1,...,ns)
24 *          xx         yy           i
25 *
26 * where
27 *
28 *          i           ns           j
29 * f (x,y,c) = c * (b(i) + sum a(i,j)*c )
30 *          i           j=1
31 *
32 * The number of species is ns = 2 * np, with the first np being
33 * prey and the last np being predators. The number np is both the
34 * number of prey and predator species. The coefficients a(i,j),
35 * b(i), d(i) are:
36 *
37 * a(i,i) = -AA  (all i)
38 * a(i,j) = -GG  (i <= np , j > np)
39 * a(i,j) = EE  (i > np, j <= np)
40 * b(i) = BB * (1 + alpha * x * y)  (i <= np)
41 * b(i) = -BB * (1 + alpha * x * y)  (i > np)
42 * d(i) = DPREY  (i <= np)
43 * d(i) = DPRED  ( i > np)
44 *
45 * The various scalar parameters are set using define's or in
46 * routine InitUserData.
47 *
48 * The boundary conditions are: normal derivative = 0, and the
49 * initial guess is constant in x and y, but the final solution
50 * is not.
51 *
52 * The PDEs are discretized by central differencing on an MX by
```

```

53 * MY mesh.
54 *
55 * The nonlinear system is solved by KINSOL using the method
56 * specified in local variable globalstrat.
57 *
58 * The preconditioner matrix is a block-diagonal matrix based on
59 * the partial derivatives of the interaction terms f only.
60 *
61 * Constraints are imposed to make all components of the solution
62 * positive.
63 * -----
64 * References:
65 *
66 * 1. Peter N. Brown and Youcef Saad,
67 *     Hybrid Krylov Methods for Nonlinear Systems of Equations
68 *     LLNL report UCRL-97645, November 1987.
69 *
70 * 2. Peter N. Brown and Alan C. Hindmarsh,
71 *     Reduced Storage Matrix Methods in Stiff ODE systems,
72 *     Lawrence Livermore National Laboratory Report UCRL-95088,
73 *     Rev. 1, June 1987, and Journal of Applied Mathematics and
74 *     Computation, Vol. 31 (May 1989), pp. 40-91. (Presents a
75 *     description of the time-dependent version of this test
76 *     problem.)
77 * -----
78 */
79
80 #include <stdio.h>
81 #include <stdlib.h>
82 #include <math.h>
83 #include "kinsol.h"          /* main KINSOL header file           */
84 #include "kinspgmr.h"        /* use KINSPGMR linear solver       */
85 #include "sundialstypes.h"   /* def's of realtype and booleantype */
86 #include "nvector_serial.h"  /* definitions of type N_Vector and access macros */
87 #include "iterative.h"        /* contains the enum for types of preconditioning */
88 #include "smalldense.h"       /* use generic DENSE solver for preconditioning */
89 #include "sundialsmath.h"     /* contains RSqrt routine           */
90
91 /* Problem Constants */
92
93 #define NUM_SPECIES      6 /* must equal 2*(number of prey or predators)
94                           number of prey = number of predators */
95
96 #define PI             RCONST(3.1415926535898) /* pi */
97
98 #define MX            8 /* MX = number of x mesh points */
99 #define MY            8 /* MY = number of y mesh points */
100 #define NSMX          (NUM_SPECIES * MX)
101 #define NEQ           (NSMX * MY) /* number of equations in the system */
102 #define AA            RCONST(1.0) /* value of coefficient AA in above eqns */
103 #define EE            RCONST(10000.) /* value of coefficient EE in above eqns */
104 #define GG            RCONST(0.5e-6) /* value of coefficient GG in above eqns */
105 #define BB            RCONST(1.0) /* value of coefficient BB in above eqns */
106 #define DPREY          RCONST(1.0) /* value of coefficient dprey above */

```

```

107 #define DPRED      RCONST(0.5)      /* value of coefficient dpred above */
108 #define ALPHA      RCONST(1.0)      /* value of coefficient alpha above */
109 #define AX         RCONST(1.0)      /* total range of x variable */
110 #define AY         RCONST(1.0)      /* total range of y variable */
111 #define FTOL       RCONST(1.e-7)     /* ftol tolerance */
112 #define STOL       RCONST(1.e-13)    /* stol tolerance */
113 #define THOUSAND   RCONST(1000.0)    /* one thousand */
114 #define ZERO        RCONST(0.)      /* 0. */
115 #define ONE         RCONST(1.0)      /* 1. */
116 #define TWO         RCONST(2.0)      /* 2. */
117 #define PREYIN      RCONST(1.0)      /* initial guess for prey concentrations. */
118 #define PREDIN      RCONST(30000.0)/* initial guess for predator concs.      */
119
120 /* User-defined vector access macro: IJ_Vptr */
121
122 /* IJ_Vptr is defined in order to translate from the underlying 3D structure
123   of the dependent variable vector to the 1D storage scheme for an N-vector.
124   IJ_Vptr(vv,i,j) returns a pointer to the location in vv corresponding to
125   indices is = 0, jx = i, jy = j.      */
126
127 #define IJ_Vptr(vv,i,j)  (&NV_Ith_S(vv, i*NUM_SPECIES + j*NSMX))
128
129 /* Type : UserData
130   contains preconditioner blocks, pivot arrays, and problem constants */
131
132 typedef struct {
133   realtype **P[MX][MY];
134   long int *pivot[MX][MY];
135   realtype **acoef, *bcoef;
136   N_Vector rates;
137   realtype *cox, *coy;
138   realtype ax, ay, dx, dy;
139   realtype uround, sqruround;
140   long int mx, my, ns, np;
141 } *UserData;
142
143 /* Functions Called by the KINSOL Solver */
144
145 static void func(N_Vector cc, N_Vector fval, void *f_data);
146
147 static int PrecSetupBD(N_Vector cc, N_Vector cscale,
148                        N_Vector fval, N_Vector fscale,
149                        void *P_data,
150                        N_Vector vtemp1, N_Vector vtemp2);
151
152 static int PrecSolveBD(N_Vector cc, N_Vector cscale,
153                        N_Vector fval, N_Vector fscale,
154                        N_Vector vv, void *P_data,
155                        N_Vector ftem);
156
157 /* Private Helper Functions */
158
159 static UserData AllocUserData(void);
160 static void InitUserData(UserData data);

```

```

161 static void FreeUserData(UserData data);
162 static void SetInitialProfiles(N_Vector cc, N_Vector sc);
163 static void PrintHeader(int globalstrategy, int maxl, int maxlrst,
164                         realtype fnormtol, realtype scsteptol);
165 static void PrintOutput(N_Vector cc);
166 static void PrintFinalStats(void *kmem);
167 static void WebRate(realtype xx, realtype yy, realtype *cxy, realtype *ratesxy,
168                      void *f_data);
169 static realtype DotProd(long int size, realtype *x1, realtype *x2);
170 static int check_flag(void *flagvalue, char *funcname, int opt);
171 /*
172 -----
173 * MAIN PROGRAM
174 -----
175 */
177
178 int main()
179 {
180     int globalstrategy;
181     realtype fnormtol, scsteptol;
182     N_Vector cc, sc, constraints;
183     UserData data;
184     int flag, maxl, maxlrst;
185     void *kmem;
186
187     cc = sc = constraints = NULL;
188     kmem = NULL;
189     data = NULL;
190
191     /* Allocate memory, and set problem data, initial values, tolerances */
192     globalstrategy = KIN_INEXACT_NEWTON;
193
194     data = AllocUserData();
195     if (check_flag((void *)data, "AllocUserData", 2)) return(1);
196     InitUserData(data);
197
198     /* Create serial vectors of length NEQ */
199     cc = N_VNew_Serial(NEQ);
200     if (check_flag((void *)cc, "N_VNew_Serial", 0)) return(1);
201     sc = N_VNew_Serial(NEQ);
202     if (check_flag((void *)sc, "N_VNew_Serial", 0)) return(1);
203     data->rates = N_VNew_Serial(NEQ);
204     if (check_flag((void *)data->rates, "N_VNew_Serial", 0)) return(1);
205
206     constraints = N_VNew_Serial(NEQ);
207     if (check_flag((void *)constraints, "N_VNew_Serial", 0)) return(1);
208     N_VConst(TWO, constraints);
209
210     SetInitialProfiles(cc, sc);
211
212     fnormtol=FTOL; scsteptol=STOL;
213
214     /* Call KINCreate/KINMalloc to initialize KINSOL:

```

```

215     nvSpec is the nvSpec pointer used in the serial version
216     A pointer to KINSOL problem memory is returned and stored in kmem. */
217     kmem = KINCreate();
218     if (check_flag((void *)kmem, "KINCreate", 0)) return(1);
219     /* Vector cc passed as template vector. */
220     flag = KINMalloc(kmem, func, cc);
221     if (check_flag(&flag, "KINMalloc", 1)) return(1);
222
223     flag = KINSetFdata(kmem, data);
224     if (check_flag(&flag, "KINSetFdata", 1)) return(1);
225     flag = KINSetConstraints(kmem, constraints);
226     if (check_flag(&flag, "KINSetConstraints", 1)) return(1);
227     flag = KINSetFuncNormTol(kmem, fnormtol);
228     if (check_flag(&flag, "KINSetFuncNormTol", 1)) return(1);
229     flag = KINSetScaledStepTol(kmem, scsteptol);
230     if (check_flag(&flag, "KINSetScaledStepTol", 1)) return(1);
231
232     /* Call KINSpgmr to specify the linear solver KINSPGMR with preconditioner
233        routines PrecSetupBD and PrecSolveBD, and the pointer to the user block data. */
234     maxl = 15;
235     maxlrst = 2;
236     flag = KINSpgmr(kmem, maxl);
237     if (check_flag(&flag, "KINSpgmr", 1)) return(1);
238
239     flag = KINSpgmrSetMaxRestarts(kmem, maxlrst);
240     if (check_flag(&flag, "KINSpgmrSetMaxRestarts", 1)) return(1);
241     flag = KINSpgmrSetPrecSetupFn(kmem, PrecSetupBD);
242     if (check_flag(&flag, "KINSpgmrSetPrecSetupFn", 1)) return(1);
243     flag = KINSpgmrSetPrecSolveFn(kmem, PrecSolveBD);
244     if (check_flag(&flag, "KINSpgmrSetPrecSolveFn", 1)) return(1);
245     flag = KINSpgmrSetPrecData(kmem, data);
246     if (check_flag(&flag, "KINSpgmrSetPrecData", 1)) return(1);
247
248     /* Print out the problem size, solution parameters, initial guess. */
249     PrintHeader(globalstrategy, maxl, maxlrst, fnormtol, scsteptol);
250
251     /* Call KINSol and print output concentration profile */
252     flag = KINSol(kmem,           /* KINSol memory block */
253                   cc,            /* initial guess on input; solution vector */
254                   globalstrategy, /* global strategy choice */
255                   sc,             /* scaling vector, for the variable cc */
256                   sc);           /* scaling vector for function values fval */
257     if (check_flag(&flag, "KINSol", 1)) return(1);
258
259     printf("\n\n\nComputed equilibrium species concentrations:\n");
260     PrintOutput(cc);
261
262     /* Print final statistics and free memory */
263     PrintFinalStats(kmem);
264
265     N_VDestroy_Serial(cc);
266     N_VDestroy_Serial(sc);
267     N_VDestroy_Serial(constraints);
268     KINFree(kmem);

```

```

269     FreeUserData(data);
270
271     return(0);
272 }
273
274 /* Readability definitions used in other routines below */
275
276 #define acoef  (data->acoef)
277 #define bcoef  (data->bcoef)
278 #define cox    (data->cox)
279 #define coy    (data->coy)
280
281 /*
282 *-----
283 * FUNCTIONS CALLED BY KINSOL
284 *-----
285 */
286
287 /*
288 * System function for predator-prey system
289 */
290
291 static void func(N_Vector cc, N_Vector fval, void *f_data)
292 {
293     realtype xx, yy, delx, dely, *cxy, *rxy, *fx, dcycli, dcyyui, dcxli, dcxri;
294     long int jx, jy, is, idyu, idyl, idxr, idxl;
295     UserData data;
296
297     data = (UserData)f_data;
298     delx = data->dx;
299     dely = data->dy;
300
301     /* Loop over all mesh points, evaluating rate array at each point*/
302     for (jy = 0; jy < MY; jy++) {
303
304         yy = dely*jy;
305
306         /* Set lower/upper index shifts, special at boundaries. */
307         idyl = (jy != 0) ? NSMX : -NSMX;
308         idyu = (jy != MY-1) ? NSMX : -NSMX;
309
310         for (jx = 0; jx < MX; jx++) {
311
312             xx = delx*jx;
313
314             /* Set left/right index shifts, special at boundaries. */
315             idxl = (jx != 0) ? NUM_SPECIES : -NUM_SPECIES;
316             idxr = (jx != MX-1) ? NUM_SPECIES : -NUM_SPECIES;
317
318             cxy = IJ_Vptr(cc, jx, jy);
319             rxy = IJ_Vptr(data->rates, jx, jy);
320             fx = IJ_Vptr(fval, jx, jy);
321
322             /* Get species interaction rate array at (xx,yy) */

```

```

323     WebRate(xx, yy, cxy, rxy, f_data);
324
325     for(is = 0; is < NUM_SPECIES; is++) {
326
327         /* Differencing in x direction */
328         dcyli = *(cxy+is) - *(cxy - idyl + is) ;
329         dcyui = *(cxy + idyu + is) - *(cxy+is);
330
331         /* Differencing in y direction */
332         dcxli = *(cxy+is) - *(cxy - idxl + is);
333         dcxri = *(cxy + idxr +is) - *(cxy+is);
334
335         /* Compute the total rate value at (xx,yy) */
336         fxy[is] = (coy)[is] * (dcyui - dcyli) +
337             (cox)[is] * (dcxri - dcxli) + rxy[is];
338
339     } /* end of is loop */
340
341 } /* end of jx loop */
342
343 } /* end of jy loop */
344 }
345
346 /*
347  * Preconditioner setup routine. Generate and preprocess P.
348  */
349
350 static int PrecSetupBD(N_Vector cc, N_Vector cscale,
351                         N_Vector fval, N_Vector fscale,
352                         void *P_data,
353                         N_Vector vtemp1, N_Vector vtemp2)
354 {
355     realtype r, r0, uround, sqruround, xx, yy, delx, dely, csave, fac;
356     realtype *cxy, *scxy, **Pxy, *ratesxy, *Pxycol, perturb_rates[NUM_SPECIES];
357     long int i, j, jx, jy, ret;
358     UserData data;
359
360     data = (UserData) P_data;
361     delx = data->dx;
362     dely = data->dy;
363
364     uround = data->uround;
365     sqruround = data->sqruround;
366     fac = N_VWL2Norm(fval, fscale);
367     r0 = THOUSAND * uround * fac * NEQ;
368     if(r0 == ZERO) r0 = ONE;
369
370     /* Loop over spatial points; get size NUM_SPECIES Jacobian block at each */
371     for (jy = 0; jy < MY; jy++) {
372         yy = jy*dely;
373
374         for (jx = 0; jx < MX; jx++) {
375             xx = jx*delx;
376             Pxy = (data->P)[jx][jy];

```

```

377     cxy = IJ_Vptr(cc,jx,jy);
378     scxy= IJ_Vptr(cscale,jx,jy);
379     ratesxy = IJ_Vptr((data->rates),jx,jy);
380
381     /* Compute difference quotients of interaction rate fn. */
382     for (j = 0; j < NUM_SPECIES; j++) {
383
384         csave = cxy[j]; /* Save the j,jx,jy element of cc */
385         r = MAX(sqrtround*ABS(csave), r0/scxy[j]);
386         cxy[j] += r; /* Perturb the j,jx,jy element of cc */
387         fac = ONE/r;
388
389         WebRate(xx, yy, cxy, perturb_rates, data);
390
391         /* Restore j,jx,jy element of cc */
392         cxy[j] = csave;
393
394         /* Load the j-th column of difference quotients */
395         Pxycol = Pxy[j];
396         for (i = 0; i < NUM_SPECIES; i++)
397             Pxycol[i] = (perturb_rates[i] - ratesxy[i]) * fac;
398
399     } /* end of j loop */
400
401     /* Do LU decomposition of size NUM_SPECIES preconditioner block */
402     ret = gefa(Pxy, NUM_SPECIES, (data->pivot)[jx][jy]);
403     if (ret != 0) return(1);
404
405 } /* end of jx loop */
406
407 } /* end of jy loop */
408
409     return(0);
410 }
411
412 /*
413  * Preconditioner solve routine
414  */
415
416
417 static int PrecSolveBD(N_Vector cc, N_Vector cscale,
418                         N_Vector fval, N_Vector fscale,
419                         N_Vector vv, void *P_data,
420                         N_Vector fitem)
421 {
422     realtype **Pxy, *vxy;
423     long int *piv, jx, jy;
424     UserData data;
425
426     data = (UserData)P_data;
427
428     for (jx=0; jx<MX; jx++) {
429         for (jy=0; jy<MY; jy++) {

```

```

431
432     /* For each (jx,jy), solve a linear system of size NUM_SPECIES.
433        vxy is the address of the corresponding portion of the vector vv;
434        Pxy is the address of the corresponding block of the matrix P;
435        piv is the address of the corresponding block of the array pivot. */
436     vxy = IJ_Vptr(vv,jx,jy);
437     Pxy = (data->P)[jx][jy];
438     piv = (data->pivot)[jx][jy];
439     gesl (Pxy, NUM_SPECIES, piv, vxy);
440
441 } /* end of jy loop */
442
443 } /* end of jx loop */
444
445 return(0);
446 }
447
448 /*
449 * Interaction rate function routine
450 */
451
452 static void WebRate(realtype xx, realtype yy, realtype *cxy, realtype *ratesxy,
453                      void *f_data)
454 {
455     long int i;
456     realtype fac;
457     UserData data;
458
459     data = (UserData)f_data;
460
461     for (i = 0; i<NUM_SPECIES; i++)
462         ratesxy[i] = DotProd(NUM_SPECIES, cxy, acoef[i]);
463
464     fac = ONE + ALPHA * xx * yy;
465
466     for (i = 0; i < NUM_SPECIES; i++)
467         ratesxy[i] = cxy[i] * ( bcoef[i] * fac + ratesxy[i] );
468 }
469
470 /*
471 * Dot product routine for realtype arrays
472 */
473
474 static realtype DotProd(long int size, realtype *x1, realtype *x2)
475 {
476     long int i;
477     realtype *xx1, *xx2, temp = ZERO;
478
479     xx1 = x1; xx2 = x2;
480     for (i = 0; i < size; i++) temp += (*xx1++) * (*xx2++);
481
482     return(temp);
483 }
484

```

```

485  /*
486  *-----
487  * PRIVATE FUNCTIONS
488  *-----
489  */
490
491 /*
492  * Allocate memory for data structure of type UserData
493  */
494
495 static UserData AllocUserData(void)
496 {
497     int jx, jy;
498     UserData data;
499
500     data = (UserData) malloc(sizeof *data);
501
502     for (jx=0; jx < MX; jx++) {
503         for (jy=0; jy < MY; jy++) {
504             (data->P)[jx][jy] = denalloc(NUM_SPECIES);
505             (data->pivot)[jx][jy] = denallocpiv(NUM_SPECIES);
506         }
507     }
508     acoef = denalloc(NUM_SPECIES);
509     bcoef = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
510     cox   = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
511     coy   = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
512
513     return(data);
514 }
515
516 /*
517  * Load problem constants in data
518  */
519
520 static void InitUserData(UserData data)
521 {
522     long int i, j, np;
523     realtype *a1,*a2, *a3, *a4, dx2, dy2;
524
525     data->mx = MX;
526     data->my = MY;
527     data->ns = NUM_SPECIES;
528     data->np = NUM_SPECIES/2;
529     data->ax = AX;
530     data->ay = AY;
531     data->dx = (data->ax)/(MX-1);
532     data->dy = (data->ay)/(MY-1);
533     data->uround = UNIT_ROUNDOFF;
534     data->sqruround = RSqrt(data->uround);
535
536     /* Set up the coefficients a and b plus others found in the equations */
537     np = data->np;
538

```

```

539     dx2=(data->dx)*(data->dx); dy2=(data->dy)*(data->dy);
540
541     for (i = 0; i < np; i++) {
542         a1= &(acoef[i][np]);
543         a2= &(acoef[i+np][0]);
544         a3= &(acoef[i][0]);
545         a4= &(acoef[i+np][np]);
546
547         /* Fill in the portion of acoef in the four quadrants, row by row */
548         for (j = 0; j < np; j++) {
549             *a1++ = -GG;
550             *a2++ = EE;
551             *a3++ = ZERO;
552             *a4++ = ZERO;
553         }
554
555         /* and then change the diagonal elements of acoef to -AA */
556         acoef[i][i]=-AA;
557         acoef[i+np][i+np] = -AA;
558
559         bcoef[i] = BB;
560         bcoef[i+np] = -BB;
561
562         cox[i]=DPREY/dx2;
563         cox[i+np]=DPRED/dx2;
564
565         coy[i]=DPREY/dy2;
566         coy[i+np]=DPRED/dy2;
567     }
568 }
569
570 /*
571  * Free data memory
572 */
573
574 static void FreeUserData(UserData data)
575 {
576     int jx, jy;
577
578     for (jx=0; jx < MX; jx++) {
579         for (jy=0; jy < MY; jy++) {
580             denfree((data->P)[jx][jy]);
581             denfreepiv((data->pivot)[jx][jy]);
582         }
583     }
584
585     denfree(acoef);
586     free(bcoef);
587     free(cox);
588     N_VDestroy_Serial(data->rates);
589     free(data);
590 }
591
592 /*

```

```

593 * Set initial conditions in cc
594 */
595
596 static void SetInitialProfiles(N_Vector cc, N_Vector sc)
597 {
598     int i, jx, jy;
599     realtype *cloc, *sloc;
600     realtype ctemp[NUM_SPECIES], stemp[NUM_SPECIES];
601
602     /* Initialize arrays ctemp and stemp used in the loading process */
603     for (i = 0; i < NUM_SPECIES/2; i++) {
604         ctemp[i] = PREYIN;
605         stemp[i] = ONE;
606     }
607     for (i = NUM_SPECIES/2; i < NUM_SPECIES; i++) {
608         ctemp[i] = PREDIN;
609         stemp[i] = RCONST(0.00001);
610     }
611
612     /* Load initial profiles into cc and sc vector from ctemp and stemp. */
613     for (jy = 0; jy < MY; jy++) {
614         for (jx = 0; jx < MX; jx++) {
615             cloc = IJ_Vptr(cc, jx, jy);
616             sloc = IJ_Vptr(sc, jx, jy);
617             for (i = 0; i < NUM_SPECIES; i++) {
618                 cloc[i] = ctemp[i];
619                 sloc[i] = stemp[i];
620             }
621         }
622     }
623 }
624
625 /*
626 * Print first lines of output (problem description)
627 */
628
629 static void PrintHeader(int globalstrategy, int maxl, int maxlrst,
630                         realtype fnormtol, realtype scsteptol)
631 {
632     printf("\nPredator-prey test problem -- KINSol (serial version)\n\n");
633     printf("Mesh dimensions = %d X %d\n", MX, MY);
634     printf("Number of species = %d\n", NUM_SPECIES);
635     printf("Total system size = %d\n\n", NEQ);
636     printf("Flag globalstrategy = %d (1 = Inex. Newton, 2 = Linesearch)\n",
637           globalstrategy);
638     printf("Linear solver is SPGMR with maxl = %d, maxlrst = %d\n",
639           maxl, maxlrst);
640     printf("Preconditioning uses interaction-only block-diagonal matrix\n");
641     printf("Positivity constraints imposed on all components \n");
642 #if defined(SUNDIALS_EXTENDED_PRECISION)
643     printf("Tolerance parameters: fnormtol = %Lg    scsteptol = %Lg\n",
644           fnormtol, scsteptol);
645 #elif defined(SUNDIALS_DOUBLE_PRECISION)
646     printf("Tolerance parameters: fnormtol = %lg    scsteptol = %lg\n",

```

```

647         fnormtol, scsteptol);
648 #else
649     printf("Tolerance parameters: fnormtol = %g    scsteptol = %g\n",
650            fnormtol, scsteptol);
651 #endif
652
653     printf("\nInitial profile of concentration\n");
654 #if defined(SUNDIALS_EXTENDED_PRECISION)
655     printf("At all mesh points: %Lg %Lg %Lg    %Lg %Lg %Lg\n",
656            PREYIN, PREYIN, PREYIN,
657            PREDIN, PREDIN, PREDIN);
658 #elif defined(SUNDIALS_DOUBLE_PRECISION)
659     printf("At all mesh points: %lg %lg %lg    %lg %lg %lg\n",
660            PREYIN, PREYIN, PREYIN,
661            PREDIN, PREDIN, PREDIN);
662 #else
663     printf("At all mesh points: %g %g %g    %g %g %g\n",
664            PREYIN, PREYIN, PREYIN,
665            PREDIN, PREDIN, PREDIN);
666 #endif
667 }
668
669 /*
670  * Print sampled values of current cc
671 */
672
673 static void PrintOutput(N_Vector cc)
674 {
675     int is, jx, jy;
676     realtype *ct;
677
678     jy = 0; jx = 0;
679     ct = IJ_Vptr(cc,jx,jy);
680     printf("\nAt bottom left:");
681
682     /* Print out lines with up to 6 values per line */
683     for (is = 0; is < NUM_SPECIES; is++){
684         if ((is%6)*6 == is) printf("\n");
685 #if defined(SUNDIALS_EXTENDED_PRECISION)
686         printf(" %Lg",ct[is]);
687 #elif defined(SUNDIALS_DOUBLE_PRECISION)
688         printf(" %lg",ct[is]);
689 #else
690         printf(" %g",ct[is]);
691 #endif
692     }
693
694     jy = MY-1; jx = MX-1;
695     ct = IJ_Vptr(cc,jx,jy);
696     printf("\n\nAt top right:");
697
698     /* Print out lines with up to 6 values per line */
699     for (is = 0; is < NUM_SPECIES; is++) {
700         if ((is%6)*6 == is) printf("\n");

```

```

701 #if defined(SUNDIALS_EXTENDED_PRECISION)
702     printf(" %Lg",ct[is]);
703 #elif defined(SUNDIALS_DOUBLE_PRECISION)
704     printf(" %lg",ct[is]);
705 #else
706     printf(" %g",ct[is]);
707 #endif
708 }
709 printf("\n\n");
710 }

711 /*
712 * Print final statistics contained in iopt
713 */
714

715 static void PrintFinalStats(void *kmem)
716 {
717     long int nni, nfe, nli, npe, nps, ncfl, nfeSG;
718     int flag;

719     flag = KINGetNumNonlinSolvIters(kmem, &nni);
720     check_flag(&flag, "KINGetNumNonlinSolvIters", 1);
721     flag = KINGetNumFuncEvals(kmem, &nfe);
722     check_flag(&flag, "KINGetNumFuncEvals", 1);
723     flag = KINSpgrmrGetNumLinIters(kmem, &nli);
724     check_flag(&flag, "KINSpgrmrGetNumLinIters", 1);
725     flag = KINSpgrmrGetNumPrecEvals(kmem, &npe);
726     check_flag(&flag, "KINSpgrmrGetNumPrecEvals", 1);
727     flag = KINSpgrmrGetNumPrecSolves(kmem, &nps);
728     check_flag(&flag, "KINSpgrmrGetNumPrecSolves", 1);
729     flag = KINSpgrmrGetNumConvFails(kmem, &ncfl);
730     check_flag(&flag, "KINSpgrmrGetNumConvFails", 1);
731     flag = KINSpgrmrGetNumFuncEvals(kmem, &nfeSG);
732     check_flag(&flag, "KINSpgrmrGetNumFuncEvals", 1);

733     printf("\nFinal Statistics.. \n\n");
734     printf("nni      = %5ld      nli      = %5ld\n", nni, nli);
735     printf("nfe      = %5ld      nfeSG    = %5ld\n", nfe, nfeSG);
736     printf("nps      = %5ld      npe      = %5ld      ncfl    = %5ld\n", nps, npe, ncfl);
737 }

738 /*
739 * Check function return value...
740 *   opt == 0 means SUNDIALS function allocates memory so check if
741 *           returned NULL pointer
742 *   opt == 1 means SUNDIALS function returns a flag so check if
743 *           flag >= 0
744 *   opt == 2 means function allocates memory so check if returned
745 *           NULL pointer
746 */
747

748 static int check_flag(void *flagvalue, char *funcname, int opt)
749 {

```

```

755     int *errflag;
756
757     /* Check if SUNDIALS function returned NULL pointer - no memory allocated */
758     if (opt == 0 && flagvalue == NULL) {
759         fprintf(stderr,
760                 "\nSUNDIALS_ERROR: %s() failed - returned NULL pointer\n\n",
761                 funcname);
762         return(1);
763     }
764
765     /* Check if flag < 0 */
766     else if (opt == 1) {
767         errflag = flagvalue;
768         if (*errflag < 0) {
769             fprintf(stderr,
770                     "\nSUNDIALS_ERROR: %s() failed with flag = %d\n\n",
771                     funcname, *errflag);
772             return(1);
773         }
774     }
775
776     /* Check if function returned NULL pointer - no memory allocated */
777     else if (opt == 2 && flagvalue == NULL) {
778         fprintf(stderr,
779                 "\nMEMORY_ERROR: %s() failed - returned NULL pointer\n\n",
780                 funcname);
781         return(1);
782     }
783
784     return(0);
785 }
```

## B Listing of kinwebbd.c

```
1  /*
2   *
3   * $Revision: 1.18 $
4   * $Date: 2004/11/15 17:28:18 $
5   *
6   * Programmer(s): Allan Taylor, Alan Hindmarsh and
7   *                  Radu Serban @ LLNL
8   *
9   * Example problem for KINSol (parallel machine case) using the BBD
10  * preconditioner.
11  *
12  * This example solves a nonlinear system that arises from a system
13  * of partial differential equations. The PDE system is a food web
14  * population model, with predator-prey interaction and diffusion on
15  * the unit square in two dimensions. The dependent variable vector
16  * is the following:
17  *
18  *      1   2           ns
19  * c = (c , c , ... , c )      (denoted by the variable cc)
20  *
21  * and the PDE's are as follows:
22  *
23  *          i      i
24  * 0 = d(i)*(c    + c    ) + f (x,y,c)  (i=1,...,ns)
25  *          xx     yy      i
26  *
27  * where
28  *
29  *          i           ns      j
30  * f (x,y,c) = c * (b(i) + sum a(i,j)*c )
31  *          i                   j=1
32  *
33  * The number of species is ns = 2 * np, with the first np being
34  * prey and the last np being predators. The number np is both the
35  * number of prey and predator species. The coefficients a(i,j),
36  * b(i), d(i) are:
37  *
38  * a(i,i) = -AA  (all i)
39  * a(i,j) = -GG  (i <= np , j > np)
40  * a(i,j) = EE  (i > np, j <= np)
41  * b(i) = BB * (1 + alpha * x * y)  (i <= np)
42  * b(i) = -BB * (1 + alpha * x * y)  (i > np)
43  * d(i) = DPREY  (i <= np)
44  * d(i) = DPRED  (i > np)
45  *
46  * The various scalar parameters are set using define's or in
47  * routine InitUserData.
48  *
49  * The boundary conditions are: normal derivative = 0, and the
50  * initial guess is constant in x and y, although the final
51  * solution is not.
52  *
```

```

53 * The PDEs are discretized by central differencing on a MX by
54 * MY mesh.
55 *
56 * The nonlinear system is solved by KINSOL using the method
57 * specified in the local variable globalstrat.
58 *
59 * The preconditioner matrix is a band-block-diagonal matrix
60 * using the KINBBDPRE module. The half-bandwidths are:
61 *
62 * ml = mu = 2*ns - 1
63 * -----
64 * References:
65 *
66 * 1. Peter N. Brown and Youcef Saad,
67 *     Hybrid Krylov Methods for Nonlinear Systems of Equations
68 *     LLNL report UCRL-97645, November 1987.
69 *
70 * 2. Peter N. Brown and Alan C. Hindmarsh,
71 *     Reduced Storage Matrix Methods in Stiff ODE systems,
72 *     Lawrence Livermore National Laboratory Report UCRL-95088,
73 *     Rev. 1, June 1987, and Journal of Applied Mathematics and
74 *     Computation, Vol. 31 (May 1989), pp. 40-91. (Presents a
75 *     description of the time-dependent version of this
76 *     test problem.)
77 * -----
78 * Run command line: mpirun -np N -machinefile machines kinwebbd
79 * where N = NPEX * NPEY is the number of processors.
80 * -----
81 */
82
83 #include <stdio.h>
84 #include <stdlib.h>
85 #include <math.h>
86 #include "sundialstypes.h"      /* def's of realtype and booleantype      */
87 #include "kinsol.h"             /* main KINSol header file                */
88 #include "iterative.h"          /* enum for types of preconditioning    */
89 #include "kinspgmr.h"           /* use KINSpgmr linear solver           */
90 #include "smalldense.h"          /* use generic DENSE solver for preconditioning*/
91 #include "nvector_parallel.h"    /* def's of type N_Vector, macro NV_DATA_P */
92 #include "sundialsmath.h"         /* contains RSqrt routine               */
93 #include "mpi.h"                /* MPI include file                     */
94 #include "kinbbdpre.h"           /* band preconditioner function prototypes */
95
96 /* Problem Constants */
97
98 #define NUM_SPECIES      6 /* must equal 2*(number of prey or predators)
99                                number of prey = number of predators */
100
101 #define PI          RCONST(3.1415926535898) /* pi */
102
103 #define NPEX        2 /* number of processors in the x-direction */
104 #define NPEY        2 /* number of processors in the y-direction */
105 #define MXSUB       10 /* number of x mesh points per subgrid */
106 #define MYSUB       10 /* number of y mesh points per subgrid */

```

```

107 #define MX          (NPEX*MXSUB) /* number of grid points in x-direction */
108 #define MY          (NPEY*MYSUB) /* number of grid points in y-direction */
109 #define NSMXSUB     (NUM_SPECIES * MXSUB)
110 #define NSMXSUB2    (NUM_SPECIES * (MXSUB+2))
111 #define NEQ          (NUM_SPECIES*MX*MY) /* number of equations in system */
112 #define AA           RCONST(1.0) /* value of coefficient AA in above eqns */
113 #define EE           RCONST(10000.) /* value of coefficient EE in above eqns */
114 #define GG           RCONST(0.5e-6) /* value of coefficient GG in above eqns */
115 #define BB           RCONST(1.0) /* value of coefficient BB in above eqns */
116 #define DPREY        RCONST(1.0) /* value of coefficient dprey above */
117 #define DPRED        RCONST(0.5) /* value of coefficient dpred above */
118 #define ALPHA         RCONST(1.0) /* value of coefficient alpha above */
119 #define AX           RCONST(1.0) /* total range of x variable */
120 #define AY           RCONST(1.0) /* total range of y variable */
121 #define FTOL          RCONST(1.e-7) /* ftol tolerance */
122 #define STOL          RCONST(1.e-13) /* stol tolerance */
123 #define THOUSAND      RCONST(1000.0) /* one thousand */
124 #define ZERO          RCONST(0.0) /* 0. */
125 #define ONE           RCONST(1.0) /* 1. */
126 #define PREYIN        RCONST(1.0) /* initial guess for prey concentrations. */
127 #define PREDIN        RCONST(30000.0)/* initial guess for predator concs. */
128
129 /* User-defined vector access macro: IJ_Vptr */
130
131 /* IJ_Vptr is defined in order to translate from the underlying 3D structure
132   of the dependent variable vector to the 1D storage scheme for an N-vector.
133   IJ_Vptr(vv,i,j) returns a pointer to the location in vv corresponding to
134   indices is = 0, jx = i, jy = j. */
135
136 #define IJ_Vptr(vv,i,j) (&NV_Ith_P(vv, i*NUM_SPECIES + j*NSMXSUB))
137
138 /* Type : UserData
139   contains preconditioner blocks, pivot arrays, and problem constants */
140
141 typedef struct {
142   realtype **acoef, *bcoef;
143   N_Vector rates;
144   realtype *cox, *coy;
145   realtype ax, ay, dx, dy;
146   long int Nlocal, mx, my, ns, np;
147   realtype cext[NUM_SPECIES * (MXSUB+2)*(MYSUB+2)];
148   long int my_pe, isubx, isuby, nsmxsub, nsmxsub2;
149   MPI_Comm comm;
150 } *UserData;
151
152 /* Function called by the KINSol Solver */
153
154 static void func(N_Vector cc, N_Vector fval, void *f_data);
155
156 static void ccomm(long int Nlocal, N_Vector cc, void *data);
157
158 static void func_local(long int Nlocal, N_Vector cc, N_Vector fval, void *f_data);
159
160 /* Private Helper Functions */

```

```

161 static UserData AllocUserData(void);
162 static void InitUserData(long int my_pe, long int Nlocal, MPI_Comm comm, UserData data);
163 static void FreeUserData(UserData data);
164 static void SetInitialProfiles(N_Vecor cc, N_Vecor sc);
165 static void PrintHeader(int globalstrategy, int maxl, int maxlrst,
166                         long int mu, long int ml,
167                         realtype fnormtol, realtype scsteptol);
168 static void PrintOutput(long int my_pe, MPI_Comm comm, N_Vecor cc);
169 static void PrintFinalStats(void *kmem);
170 static void WebRate(realtype xx, realtype yy, realtype *cxy, realtype *ratesxy,
171                     void *f_data);
172 static realtype DotProd(long int size, realtype *x1, realtype *x2);
173 static void BSend(MPI_Comm comm, long int my_pe, long int isubx,
174                   long int isuby, long int dsizex, long int dsizey,
175                   realtype *cdata);
176 static void BRecvPost(MPI_Comm comm, MPI_Request request[], long int my_pe,
177                      long int isubx, long int isuby,
178                      long int dsizex, long int dsizey,
179                      realtype *cext, realtype *buffer);
180 static void BRecvWait(MPI_Request request[], long int isubx,
181                      long int isuby, long int dsizex, realtype *cext,
182                      realtype *buffer);
183 static int check_flag(void *flagvalue, char *funcname, int opt, int id);
184
185 /*
186 -----
187 * MAIN PROGRAM
188 -----
189 */
190
191
192 int main(int argc, char *argv[])
193 {
194     MPI_Comm comm;
195     void *kmem, *pdata;
196     UserData data;
197     N_Vecor cc, sc, constraints;
198     int globalstrategy;
199     long int Nlocal;
200     realtype fnormtol, scsteptol, dq_rel_uu;
201     int flag, maxl, maxlrst;
202     long int mu, ml;
203     int my_pe, npes, npelast = NPEX*NPEY-1;
204
205     data = NULL;
206     kmem = pdata = NULL;
207     cc = sc = constraints = NULL;
208
209     /* Get processor number and total number of pe's */
210     MPI_Init(&argc, &argv);
211     comm = MPI_COMM_WORLD;
212     MPI_Comm_size(comm, &npes);
213     MPI_Comm_rank(comm, &my_pe);
214

```

```

215 if (npes != NPEX*NPEY) {
216     if (my_pe == 0)
217         printf("\nMPI_ERROR(0): npes=%d is not equal to NPEX*NPEY=%d\n", npes, NPEX*NPEY);
218     return(1);
219 }
220
221 /* Allocate memory, and set problem data, initial values, tolerances */
222
223 /* Set local length */
224 Nlocal = NUM_SPECIES*MXSUB*MYSUB;
225
226 /* Allocate and initialize user data block */
227 data = AllocUserData();
228 if (check_flag((void *)data, "AllocUserData", 2, my_pe)) MPI_Abort(comm, 1);
229 InitUserData(my_pe, Nlocal, comm, data);
230
231 /* Choose global strategy */
232 globalstrategy = KIN_INEXACT_NEWTON;
233
234 /* Allocate and initialize vectors */
235 cc = N_VNew_Parallel(comm, Nlocal, NEQ);
236 if (check_flag((void *)cc, "N_VNew_Parallel", 0, my_pe)) MPI_Abort(comm, 1);
237 sc = N_VNew_Parallel(comm, Nlocal, NEQ);
238 if (check_flag((void *)sc, "N_VNew_Parallel", 0, my_pe)) MPI_Abort(comm, 1);
239 data->rates = N_VNew_Parallel(comm, Nlocal, NEQ);
240 if (check_flag((void *)data->rates, "N_VNew_Parallel", 0, my_pe)) MPI_Abort(comm, 1);
241 constraints = N_VNew_Parallel(comm, Nlocal, NEQ);
242 if (check_flag((void *)constraints, "N_VNew_Parallel", 0, my_pe)) MPI_Abort(comm, 1);
243 N_VConst(ZERO, constraints);
244
245 SetInitialProfiles(cc, sc);
246
247 fnormtol = FTOL; scsteptol = STOL;
248
249 /* Call KINCreate/KINMalloc to initialize KINSOL:
250    nvSpec points to machine environment data
251    A pointer to KINSOL problem memory is returned and stored in kmem. */
252 kmem = KINCreate();
253 if (check_flag((void *)kmem, "KINCreate", 0, my_pe)) MPI_Abort(comm, 1);
254
255 /* Vector cc passed as template vector. */
256 flag = KINMalloc(kmem, func, cc);
257 if (check_flag(&flag, "KINMalloc", 1, my_pe)) MPI_Abort(comm, 1);
258
259 flag = KINSetFdata(kmem, data);
260 if (check_flag(&flag, "KINSetFdata", 1, my_pe)) MPI_Abort(comm, 1);
261
262 flag = KINSetConstraints(kmem, constraints);
263 if (check_flag(&flag, "KINSetConstraints", 1, my_pe)) MPI_Abort(comm, 1);
264
265 flag = KINSetFuncNormTol(kmem, fnormtol);
266 if (check_flag(&flag, "KINSetFuncNormTol", 1, my_pe)) MPI_Abort(comm, 1);
267
268 flag = KINSetScaledStepTol(kmem, scsteptol);

```

```

269 if (check_flag(&flag, "KINSetScaledStepTol", 1, my_pe)) MPI_Abort(comm, 1);
270
271 /* Call KINBBDPrecAlloc to initialize and allocate memory for the
272    band-block-diagonal preconditioner, and specify the local and
273    communication functions func_local and gcomm=NULL (all communication
274    needed for the func_local is already done in func). */
275 dq_rel_uu = ZERO;
276 mu = ml = 2*NUM_SPECIES - 1;
277
278 pdata = KINBBDPrecAlloc(kmem, Nlocal, mu, ml, dq_rel_uu, func_local, NULL);
279 if (check_flag((void *)pdata, "KINBBDPrecAlloc", 0, my_pe))
280     MPI_Abort(comm, 1);
281
282 /* Call KINBBDSpgrmr to specify the linear solver KINSPGMR
283    with preconditioner KINBBDPRE */
284 maxl = 20; maxlrst = 2;
285 flag = KINBBDSpgrmr(kmem, maxl, pdata);
286 if (check_flag(&flag, "KINBBDSpgrmr", 1, my_pe))
287     MPI_Abort(comm, 1);
288
289 flag = KINSpgmrSetMaxRestarts(kmem, maxlrst);
290 if (check_flag(&flag, "KINSpgmrSetMaxRestarts", 1, my_pe))
291     MPI_Abort(comm, 1);
292
293 /* Print out the problem size, solution parameters, initial guess. */
294 if (my_pe == 0)
295     PrintHeader(globalstrategy, maxl, maxlrst, mu, ml, fnormtol, scsteptol);
296
297 /* call KINSol and print output concentration profile */
298 flag = KINSol(kmem,           /* KINSol memory block */
299                 cc,             /* initial guesss on input; solution vector */
300                 globalstrategy, /* global stragegy choice */
301                 sc,             /* scaling vector, for the variable cc */
302                 sc);           /* scaling vector for function values fval */
303 if (check_flag(&flag, "KINSol", 1, my_pe)) MPI_Abort(comm, 1);
304
305 if (my_pe == 0) printf("\n\n\nComputed equilibrium species concentrations:\n");
306 if (my_pe == 0 || my_pe==npelast) PrintOutput(my_pe, comm, cc);
307
308 /* Print final statistics and free memory */
309 if (my_pe == 0)
310     PrintFinalStats(kmem);
311
312 N_VDestroy_Parallel(cc);
313 N_VDestroy_Parallel(sc);
314 N_VDestroy_Parallel(constraints);
315 KINBBDPrecFree(pdata);
316 KINFree(kmem);
317 FreeUserData(data);
318
319 MPI_Finalize();
320
321 return(0);
322 }
```

```

323
324 /* Readability definitions used in other routines below */
325
326 #define acoef (data->acoef)
327 #define bcoef (data->bcoef)
328 #define cox (data->cox)
329 #define coy (data->coy)
330
331 /*
332 *-----
333 * FUNCTIONS CALLED BY KINSOL
334 *-----
335 */
336
337 /*
338 * ccomm routine. This routine performs all communication
339 * between processors of data needed to calculate f.
340 */
341
342 static void ccomm(long int Nlocal, N_Vector cc, void *userdata)
343 {
344
345     realtype *cdata, *cext, buffer[2*NUM_SPECIES*MYSUB];
346     UserData data;
347     MPI_Comm comm;
348     long int my_pe, isubx, isuby, nsmxsub, nsmysub;
349     MPI_Request request[4];
350
351     /* Get comm, my_pe, subgrid indices, data sizes, extended array cext */
352     data = (UserData) userdata;
353     comm = data->comm; my_pe = data->my_pe;
354     isubx = data->isubx; isuby = data->isuby;
355     nsmxsub = data->nsmxsub;
356     nsmysub = NUM_SPECIES*MYSUB;
357     cext = data->cext;
358
359     cdata = NV_DATA_P(cc);
360
361     /* Start receiving boundary data from neighboring PEs */
362     BRecvPost(comm, request, my_pe, isubx, isuby, nsmxsub, nsmysub, cext, buffer);
363
364     /* Send data from boundary of local grid to neighboring PEs */
365     BSend(comm, my_pe, isubx, isuby, nsmxsub, nsmysub, cdata);
366
367     /* Finish receiving boundary data from neighboring PEs */
368     BRecvWait(request, isubx, isuby, nsmxsub, cext, buffer);
369 }
370
371 /*
372 * System function for predator-prey system - calculation part
373 */
374
375
376 static void func_local(long int Nlocal, N_Vector cc, N_Vector fval, void *f_data)

```

```

377  {
378      realtype xx, yy, *cxy, *rxy, *fxy, dcydi, dcyui, dcxli, dcxri;
379      realtype *cext, dely, delx, *cdata;
380      long int i, jx, jy, is, ly;
381      long int isubx, isuby, nsmxsub, nsmxsub2;
382      long int shifty, offsetc, offsetce, offsetcl, offsetcr, offsetcd, offsetcu;
383      UserData data;
384
385      data = (UserData)f_data;
386      cdata = NV_DATA_P(cc);
387
388      /* Get subgrid indices, data sizes, extended work array cext */
389      isubx = data->isubx;    isuby = data->isuby;
390      nsmxsub = data->nsmxsub; nsmxsub2 = data->nsmxsub2;
391      cext = data->cext;
392
393      /* Copy local segment of cc vector into the working extended array cext */
394      offsetc = 0;
395      offsetce = nsmxsub2 + NUM_SPECIES;
396      for (ly = 0; ly < MYSUB; ly++) {
397          for (i = 0; i < nsmxsub; i++) cext[offsetce+i] = cdata[offsetc+i];
398          offsetc = offsetc + nsmxsub;
399          offsetce = offsetce + nsmxsub2;
400      }
401
402      /* To facilitate homogeneous Neumann boundary conditions, when this is a
403         boundary PE, copy data from the first interior mesh line of cc to cext */
404
405      /* If isuby = 0, copy x-line 2 of cc to cext */
406      if (isuby == 0) {
407          for (i = 0; i < nsmxsub; i++) cext[NUM_SPECIES+i] = cdata[nsmxsub+i];
408      }
409
410      /* If isuby = NPEY-1, copy x-line MYSUB-1 of cc to cext */
411      if (isuby == NPEY-1) {
412          offsetc = (MYSUB-2)*nsmxsub;
413          offsetce = (MYSUB+1)*nsmxsub2 + NUM_SPECIES;
414          for (i = 0; i < nsmxsub; i++) cext[offsetce+i] = cdata[offsetc+i];
415      }
416
417      /* If isubx = 0, copy y-line 2 of cc to cext */
418      if (isubx == 0) {
419          for (ly = 0; ly < MYSUB; ly++) {
420              offsetc = ly*nsmxsub + NUM_SPECIES;
421              offsetce = (ly+1)*nsmxsub2;
422              for (i = 0; i < NUM_SPECIES; i++) cext[offsetce+i] = cdata[offsetc+i];
423          }
424      }
425
426      /* If isubx = NPEX-1, copy y-line MXSUB-1 of cc to cext */
427      if (isubx == NPEX-1) {
428          for (ly = 0; ly < MYSUB; ly++) {
429              offsetc = (ly+1)*nsmxsub - 2*NUM_SPECIES;
430              offsetce = (ly+2)*nsmxsub2 - NUM_SPECIES;

```

```

431     for (i = 0; i < NUM_SPECIES; i++) cext[offsetce+i] = cdata[offsetc+i];
432   }
433 }
434
435 /* Loop over all mesh points, evaluating rate arra at each point */
436 delx = data->dx;
437 dely = data->dy;
438 shifty = (MXSUB+2)*NUM_SPECIES;
439
440 for (jy = 0; jy < MYSUB; jy++) {
441
442   yy = dely*(jy + isuby * MYSUB);
443
444   for (jx = 0; jx < MXSUB; jx++) {
445
446     xx = delx * (jx + isubx * MXSUB);
447     cxy = IJ_Vptr(cc,jx,jy);
448     rxy = IJ_Vptr(data->rates,jx,jy);
449     fxy = IJ_Vptr(fval,jx,jy);
450
451     WebRate(xx, yy, cxy, rxy, f_data);
452
453     offsetc = (jx+1)*NUM_SPECIES + (jy+1)*NSMXSUB2;
454     offsetcd = offsetc - shifty;
455     offsetcu = offsetc + shifty;
456     offsetcl = offsetc - NUM_SPECIES;
457     offsetcr = offsetc + NUM_SPECIES;
458
459     for (is = 0; is < NUM_SPECIES; is++) {
460
461       /* differencing in x */
462       dcydi = cext[offsetc+is] - cext[offsetcd+is];
463       dcyui = cext[offsetcu+is] - cext[offsetc+is];
464
465       /* differencing in y */
466       dcxli = cext[offsetc+is] - cext[offsetcl+is];
467       dcxri = cext[offsetcr+is] - cext[offsetc+is];
468
469       /* compute the value at xx , yy */
470       fxy[is] = (coy)[is] * (dcyui - dcydi) +
471                 (cox)[is] * (dcxri - dcxli) + rxy[is];
472
473     } /* end of is loop */
474
475   } /* end of jx loop */
476
477 } /* end of jy loop */
478 }
479
480 /*
481  * System function routine. Evaluate f(cc). First call ccomm to do
482  * communication of subgrid boundary data into cext. Then calculate f
483  * by a call to func_local.
484 */

```

```

485
486 static void func(N_Vector cc, N_Vector fval, void *f_data)
487 {
488     UserData data;
489
490     data = (UserData) f_data;
491
492     /* Call ccomm to do inter-processor communicaiton */
493     ccomm(data->Nlocal, cc, data);
494
495     /* Call func_local to calculate all right-hand sides */
496     func_local(data->Nlocal, cc, fval, data);
497 }
498
499 /*
500  * Interaction rate function routine
501 */
502
503 static void WebRate(realtype xx, realtype yy, realtype *cxy, realtype *ratesxy,
504                      void *f_data)
505 {
506     long int i;
507     realtype fac;
508     UserData data;
509
510     data = (UserData)f_data;
511
512     for (i = 0; i<NUM_SPECIES; i++)
513         ratesxy[i] = DotProd(NUM_SPECIES, cxy, acoef[i]);
514
515     fac = ONE + ALPHA * xx * yy;
516
517     for (i = 0; i < NUM_SPECIES; i++)
518         ratesxy[i] = cxy[i] * ( bcoef[i] * fac + ratesxy[i] );
519 }
520
521 /*
522  * Dot product routine for realtype arrays
523 */
524
525 static realtype DotProd(long int size, realtype *x1, realtype *x2)
526 {
527     long int i;
528     realtype *xx1, *xx2, temp = ZERO;
529
530     xx1 = x1; xx2 = x2;
531     for (i = 0; i < size; i++) temp += (*xx1++) * (*xx2++);
532
533     return(temp);
534 }
535
536 /*
537  *-----
538  * PRIVATE FUNCTIONS

```

```

539  *-
540  */
541
542 /* 
543  * Allocate memory for data structure of type UserData
544  */
545
546 static UserData AllocUserData(void)
547 {
548     UserData data;
549
550     data = (UserData) malloc(sizeof *data);
551
552     acoef = denalloc(NUM_SPECIES);
553     bcoef = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
554     cox   = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
555     coy   = (realtype *)malloc(NUM_SPECIES * sizeof(realtype));
556
557     return(data);
558 }
559
560 /*
561  * Load problem constants in data
562  */
563
564 static void InitUserData(long int my_pe, long int Nlocal, MPI_Comm comm, UserData data)
565 {
566     long int i, j, np;
567     realtype *a1,*a2, *a3, *a4, dx2, dy2;
568
569     data->mx = MX;
570     data->my = MY;
571     data->ns = NUM_SPECIES;
572     data->np = NUM_SPECIES/2;
573     data->ax = AX;
574     data->ay = AY;
575     data->dx = (data->ax)/(MX-1);
576     data->dy = (data->ay)/(MY-1);
577     data->my_pe = my_pe;
578     data->Nlocal = Nlocal;
579     data->comm = comm;
580     data->isuby = my_pe/NPEX;
581     data->isubx = my_pe - data->isuby*NPEX;
582     data->nsmxsub = NUM_SPECIES * MXSUB;
583     data->nsmxsub2 = NUM_SPECIES * (MXSUB+2);
584
585     /* Set up the coefficients a and b plus others found in the equations */
586     np = data->np;
587
588     dx2=(data->dx)*(data->dx); dy2=(data->dy)*(data->dy);
589
590     for (i = 0; i < np; i++) {
591         a1= &(acoef[i][np]);
592         a2= &(acoef[i+np][0]);

```

```

593     a3= &(acoef[i][0]);
594     a4= &(acoef[i+np][np]);
595
596     /* Fill in the portion of acoef in the four quadrants, row by row */
597     for (j = 0; j < np; j++) {
598         *a1++ = -GG;
599         *a2++ = EE;
600         *a3++ = ZERO;
601         *a4++ = ZERO;
602     }
603
604     /* and then change the diagonal elements of acoef to -AA */
605     acoef[i][i]=-AA;
606     acoef[i+np][i+np] = -AA;
607
608     bcoef[i] = BB;
609     bcoef[i+np] = -BB;
610
611     cox[i]=DPREY/dx2;
612     cox[i+np]=DPRED/dx2;
613
614     coy[i]=DPREY/dy2;
615     coy[i+np]=DPRED/dy2;
616 }
617 }
618 /*
619  * Free data memory
620 */
621
622 static void FreeUserData(UserData data)
623 {
624
625     denfree(acoef);
626     free(bcoef);
627     free(cox); free(coy);
628     N_VDestroy_Parallel(data->rates);
629
630     free(data);
631
632 }
633 /*
634  * Set initial conditions in cc
635 */
636
637 static void SetInitialProfiles(N_Vector cc, N_Vector sc)
638 {
639     int i, jx, jy;
640     realtype *cloc, *sloc;
641     realtype ctemp[NUM_SPECIES], stemp[NUM_SPECIES];
642
643     /* Initialize arrays ctemp and stemp used in the loading process */
644     for (i = 0; i < NUM_SPECIES/2; i++) {

```

```

647     ctemp[i] = PREYIN;
648     stemp[i] = ONE;
649 }
650 for (i = NUM_SPECIES/2; i < NUM_SPECIES; i++) {
651     ctemp[i] = PREDIN;
652     stemp[i] = RCONST(0.00001);
653 }
654
655 /* Load initial profiles into cc and sc vector from ctemp and stemp. */
656 for (jy = 0; jy < MYSUB; jy++) {
657     for (jx=0; jx < MXSUB; jx++) {
658         cloc = IJ_Vptr(cc,jx,jy);
659         sloc = IJ_Vptr(sc,jx,jy);
660         for (i = 0; i < NUM_SPECIES; i++){
661             cloc[i] = ctemp[i];
662             sloc[i] = stemp[i];
663         }
664     }
665 }
666
667 }
668
669 /*
670 * Print first lines of output (problem description)
671 */
672
673 static void PrintHeader(int globalstrategy, int maxl, int maxlrst,
674                         long int mu, long int ml,
675                         realtype fnormtol, realtype scsteptol)
676 {
677     printf("\nPredator-prey test problem-- KINSol (parallel-BBD version)\n\n");
678
679     printf("Mesh dimensions = %d X %d\n", MX, MY);
680     printf("Number of species = %d\n", NUM_SPECIES);
681     printf("Total system size = %d\n\n", NEQ);
682     printf("Subgrid dimensions = %d X %d\n", MXSUB, MYSUB);
683     printf("Processor array is %d X %d\n", NPEX, NPEY);
684     printf("Flag globalstrategy = %d (1 = Inex. Newton, 2 = Linesearch)\n",
685            globalstrategy);
686     printf("Linear solver is SPGMR with maxl = %d, maxlrst = %d\n",
687            maxl, maxlrst);
688     printf("Preconditioning uses band-block-diagonal matrix from KINBBDPRE\n");
689     printf(" with matrix half-bandwidths ml, mu = %ld %ld\n", ml, mu);
690 #if defined(SUNDIALS_EXTENDED_PRECISION)
691     printf("Tolerance parameters: fnormtol = %Lg    scsteptol = %Lg\n",
692           fnormtol, scsteptol);
693 #elif defined(SUNDIALS_DOUBLE_PRECISION)
694     printf("Tolerance parameters: fnormtol = %lg    scsteptol = %lg\n",
695           fnormtol, scsteptol);
696 #else
697     printf("Tolerance parameters: fnormtol = %g    scsteptol = %g\n",
698           fnormtol, scsteptol);
699 #endif
700

```

```

701     printf("\nInitial profile of concentration\n");
702 #if defined(SUNDIALS_EXTENDED_PRECISION)
703     printf("At all mesh points: %Lg %Lg %Lg    %Lg %Lg %Lg\n", PREYIN,PREYIN,PREYIN,
704           PREDIN,PREDIN,PREDIN);
705 #elif defined(SUNDIALS_DOUBLE_PRECISION)
706     printf("At all mesh points: %lg %lg %lg    %lg %lg %lg\n", PREYIN,PREYIN,PREYIN,
707           PREDIN,PREDIN,PREDIN);
708 #else
709     printf("At all mesh points: %g %g %g    %g %g %g\n", PREYIN,PREYIN,PREYIN,
710           PREDIN,PREDIN,PREDIN);
711 #endif
712 }
713
714 /*
715  * Print sample of current cc values
716 */
717
718 static void PrintOutput(long int my_pe, MPI_Comm comm, N_Vecor cc)
719 {
720     int is, i0, npelast;
721     realtype *ct, tempc[NUM_SPECIES];
722     MPI_Status status;
723
724     npelast = NPEX*NPEY - 1;
725
726     ct = NV_DATA_P(cc);
727
728     /* Send the cc values (for all species) at the top right mesh point to PE 0 */
729     if (my_pe == npelast) {
730         i0 = NUM_SPECIES*(MXSUB*MYSUB-1);
731         if (npelast!=0)
732             MPI_Send(&ct[i0],NUM_SPECIES,PVEC_REAL_MPI_TYPE,0,0,comm);
733         else /* single processor case */
734             for (is = 0; is < NUM_SPECIES; is++) tempc[is]=ct[i0+is];
735     }
736
737     /* On PE 0, receive the cc values at top right, then print performance data
738      and sampled solution values */
739     if (my_pe == 0) {
740
741         if (npelast != 0)
742             MPI_Recv(&tempc[0],NUM_SPECIES,PVEC_REAL_MPI_TYPE,npelast,0,comm,&status);
743
744         printf("\nAt bottom left:");
745         for (is = 0; is < NUM_SPECIES; is++){
746             if ((is%6)*6== is) printf("\n");
747 #if defined(SUNDIALS_EXTENDED_PRECISION)
748             printf(" %Lg",ct[is]);
749 #elif defined(SUNDIALS_DOUBLE_PRECISION)
750             printf(" %lg",ct[is]);
751 #else
752             printf(" %g",ct[is]);
753 #endif
754         }

```

```

755     printf("\n\nAt top right:");
756     for (is = 0; is < NUM_SPECIES; is++) {
757         if ((is%6)*6 == is) printf("\n");
758 #if defined(SUNDIALS_EXTENDED_PRECISION)
759         printf(" %Lg",tempc[is]);
760 #elif defined(SUNDIALS_DOUBLE_PRECISION)
761         printf(" %lg",tempc[is]);
762 #else
763         printf(" %g",tempc[is]);
764 #endif
765     }
766 }
767     printf("\n\n");
768 }
769 }
770 */
771 /*
772 * Print final statistics contained in iopt
773 */
774
775 static void PrintFinalStats(void *kmem)
776 {
777     long int nni, nfe, nli, npe, nps, ncfl, nfeSG;
778     int flag;
779
780     flag = KINGetNumNonlinSolvIters(kmem, &nni);
781     check_flag(&flag, "KINGetNumNonlinSolvIters", 1, 0);
782     flag = KINGetNumFuncEvals(kmem, &nfe);
783     check_flag(&flag, "KINGetNumFuncEvals", 1, 0);
784     flag = KINSpgmrGetNumLinIters(kmem, &nli);
785     check_flag(&flag, "KINSpgmrGetNumLinIters", 1, 0);
786     flag = KINSpgmrGetNumPrecEvals(kmem, &npe);
787     check_flag(&flag, "KINSpgmrGetNumPrecEvals", 1, 0);
788     flag = KINSpgmrGetNumPrecSolves(kmem, &nps);
789     check_flag(&flag, "KINSpgmrGetNumPrecSolves", 1, 0);
790     flag = KINSpgmrGetNumConvFails(kmem, &ncfl);
791     check_flag(&flag, "KINSpgmrGetNumConvFails", 1, 0);
792     flag = KINSpgmrGetNumFuncEvals(kmem, &nfeSG);
793     check_flag(&flag, "KINSpgmrGetNumFuncEvals", 1, 0);
794
795     printf("\nFinal Statistics.. \n\n");
796     printf("nni      = %5ld      nli      = %5ld\n", nni, nli);
797     printf("nfe      = %5ld      nfeSG    = %5ld\n", nfe, nfeSG);
798     printf("nps      = %5ld      npe      = %5ld      ncfl    = %5ld\n", nps, npe, ncfl);
799 }
800 */
801 /*
802 * Routine to send boundary data to neighboring PEs
803 */
804
805
806 static void BSend(MPI_Comm comm, long int my_pe,
807                     long int isubx, long int isuby,
808                     long int dsizex, long int dsizey, realtype *cdata)

```

```

809  {
810      int i, ly;
811      long int offsetc, offsetbuf;
812      realtype bufleft[NUM_SPECIES*MYSUB], bufright[NUM_SPECIES*MYSUB];
813
814      /* If isuby > 0, send data from bottom x-line of u */
815      if (isuby != 0)
816          MPI_Send(&cdata[0], dsizex, PVEC_REAL_MPI_TYPE, my_pe-NPEX, 0, comm);
817
818      /* If isuby < NPEY-1, send data from top x-line of u */
819      if (isuby != NPEY-1) {
820          offsetc = (MYSUB-1)*dsizex;
821          MPI_Send(&cdata[offsetc], dsizex, PVEC_REAL_MPI_TYPE, my_pe+NPEX, 0, comm);
822      }
823
824      /* If isubx > 0, send data from left y-line of u (via bufleft) */
825      if (isubx != 0) {
826          for (ly = 0; ly < MYSUB; ly++) {
827              offsetbuf = ly*NUM_SPECIES;
828              offsetc = ly*dsizex;
829              for (i = 0; i < NUM_SPECIES; i++)
830                  bufleft[offsetbuf+i] = cdata[offsetc+i];
831          }
832          MPI_Send(&bufleft[0], dsizey, PVEC_REAL_MPI_TYPE, my_pe-1, 0, comm);
833      }
834
835      /* If isubx < NPEX-1, send data from right y-line of u (via bufright) */
836      if (isubx != NPEX-1) {
837          for (ly = 0; ly < MYSUB; ly++) {
838              offsetbuf = ly*NUM_SPECIES;
839              offsetc = offsetbuf*MXSUB + (MXSUB-1)*NUM_SPECIES;
840              for (i = 0; i < NUM_SPECIES; i++)
841                  bufright[offsetbuf+i] = cdata[offsetc+i];
842          }
843          MPI_Send(&bufright[0], dsizey, PVEC_REAL_MPI_TYPE, my_pe+1, 0, comm);
844      }
845  }
846
847  /*
848   * Routine to start receiving boundary data from neighboring PEs.
849   * Notes:
850   * 1) buffer should be able to hold 2*NUM_SPECIES*MYSUB realtype entries,
851   *     should be passed to both the BRecvPost and BRecvWait functions, and
852   *     should not be manipulated between the two calls.
853   * 2) request should have 4 entries, and should be passed in both calls also.
854   */
855
856  static void BRecvPost(MPI_Comm comm, MPI_Request request[], long int my_pe,
857                        long int isubx, long int isuby,
858                        long int dsizex, long int dsizey,
859                        realtype *cext, realtype *buffer)
860  {
861      long int offsetce;
862

```

```

863 /* Have bufleft and bufright use the same buffer */
864 realtype *bufleft = buffer, *bufright = buffer+NUM_SPECIES*MYSUB;
865
866 /* If isuby > 0, receive data for bottom x-line of cext */
867 if (isuby != 0)
868     MPI_Irecv(&cext[NUM_SPECIES], dsizex, PVEC_REAL_MPI_TYPE,
869                 my_pe-NPEX, 0, comm, &request[0]);
870
871 /* If isuby < NPEY-1, receive data for top x-line of cext */
872 if (isuby != NPEY-1) {
873     offsetce = NUM_SPECIES*(1 + (MYSUB+1)*(MXSUB+2));
874     MPI_Irecv(&cext[offsetce], dsizex, PVEC_REAL_MPI_TYPE,
875                 my_pe+NPEX, 0, comm, &request[1]);
876 }
877
878 /* If isubx > 0, receive data for left y-line of cext (via bufleft) */
879 if (isubx != 0) {
880     MPI_Irecv(&bufleft[0], dsizey, PVEC_REAL_MPI_TYPE,
881                 my_pe-1, 0, comm, &request[2]);
882 }
883
884 /* If isubx < NPEX-1, receive data for right y-line of cext (via bufright) */
885 if (isubx != NPEX-1) {
886     MPI_Irecv(&bufright[0], dsizey, PVEC_REAL_MPI_TYPE,
887                 my_pe+1, 0, comm, &request[3]);
888 }
889 }
890
891 /*
892 * Routine to finish receiving boundary data from neighboring PEs.
893 * Notes:
894 * 1) buffer should be able to hold 2*NUM_SPECIES*MYSUB realtype entries,
895 *    should be passed to both the BRecvPost and BRecvWait functions, and
896 *    should not be manipulated between the two calls.
897 * 2) request should have 4 entries, and should be passed in both calls also.
898 */
899
900 static void BRecvWait(MPI_Request request[], long int isubx,
901                       long int isuby, long int dsizex, realtype *cext,
902                       realtype *buffer)
903 {
904     int i, ly;
905     long int dsizex2, offsetce, offsetbuf;
906     realtype *bufleft = buffer, *bufright = buffer+NUM_SPECIES*MYSUB;
907     MPI_Status status;
908
909     dsizex2 = dsizex + 2*NUM_SPECIES;
910
911     /* If isuby > 0, receive data for bottom x-line of cext */
912     if (isuby != 0)
913         MPI_Wait(&request[0], &status);
914
915     /* If isuby < NPEY-1, receive data for top x-line of cext */
916     if (isuby != NPEY-1)

```

```

917 MPI_Wait(&request[1],&status);
918
919 /* If isubx > 0, receive data for left y-line of cext (via bufleft) */
920 if (isubx != 0) {
921     MPI_Wait(&request[2],&status);
922
923     /* Copy the buffer to cext */
924     for (ly = 0; ly < MYSUB; ly++) {
925         offsetbuf = ly*NUM_SPECIES;
926         offsetce = (ly+1)*dsizex2;
927         for (i = 0; i < NUM_SPECIES; i++)
928             cext[offsetce+i] = bufleft[offsetbuf+i];
929     }
930 }
931
932 /* If isubx < NPEX-1, receive data for right y-line of cext (via bufright) */
933 if (isubx != NPEX-1) {
934     MPI_Wait(&request[3],&status);
935
936     /* Copy the buffer to cext */
937     for (ly = 0; ly < MYSUB; ly++) {
938         offsetbuf = ly*NUM_SPECIES;
939         offsetce = (ly+2)*dsizex2 - NUM_SPECIES;
940         for (i = 0; i < NUM_SPECIES; i++)
941             cext[offsetce+i] = bufright[offsetbuf+i];
942     }
943 }
944 /*
945 * Check function return value...
946 *   opt == 0 means SUNDIALS function allocates memory so check if
947 *           returned NULL pointer
948 *   opt == 1 means SUNDIALS function returns a flag so check if
949 *           flag >= 0
950 *   opt == 2 means function allocates memory so check if returned
951 *           NULL pointer
952 */
953
954
955 static int check_flag(void *flagvalue, char *funcname, int opt, int id)
956 {
957     int *errflag;
958
959     /* Check if SUNDIALS function returned NULL pointer - no memory allocated */
960     if (opt == 0 && flagvalue == NULL) {
961         fprintf(stderr,
962                 "\nSUNDIALS_ERROR(%d): %s() failed - returned NULL pointer\n\n",
963                 id, funcname);
964         return(1);
965     }
966
967     /* Check if flag < 0 */
968     else if (opt == 1) {
969         errflag = flagvalue;
970         if (*errflag < 0) {

```

```
971     fprintf(stderr,
972             "\nSUNDIALS_ERROR(%d): %s() failed with flag = %d\n\n",
973             id, funcname, *errflag);
974     return(1);
975 }
976 }
977
978 /* Check if function returned NULL pointer - no memory allocated */
979 else if (opt == 2 && flagvalue == NULL) {
980     fprintf(stderr,
981             "\nMEMORY_ERROR(%d): %s() failed - returned NULL pointer\n\n",
982             id, funcname);
983     return(1);
984 }
985
986 return(0);
987 }
```

## C Listing of kindiagsf.f

```
1      program kindiagsf
2      C -----
3      C $Revision: 1.13 $
4      C $Date: 2004/10/15 00:03:54 $
5      C -----
6      C Programmer(s): Allan Taylor, Alan Hindmarsh and
7      C Radu Serban @ LLNL
8      C -----
9      C Simple diagonal test with Fortran interface, using user-supplied
10     C preconditioner setup and solve routines (supplied in Fortran).
11     C
12     C This example does a basic test of the solver by solving the
13     C system:
14     C           f(u) = 0   for
15     C           f(u) = u(i)^2 - i^2
16     C
17     C No scaling is done.
18     C An approximate diagonal preconditioner is used.
19     C
20     C Execution command: kindiagsf
21     C -----
22     C
23     implicit none
24
25     integer ier, globalstrat, inopt, maxl, maxlrst
26     integer*4 PROBSIZE
27     parameter(PROBSIZE=128)
28     integer*4 neq, i, msbpre
29     integer*4 iopt(40)
30     double precision pp, fnormtol, scsteptol
31     double precision ropt(40), uu(PROBSIZE), scale(PROBSIZE)
32     double precision constr(PROBSIZE)
33
34     common /pcom/ pp(PROBSIZE)
35     common /psize/ neq
36
37     neq = PROBSIZE
38     globalstrat = 1
39     fnormtol = 1.0d-5
40     scsteptol = 1.0d-4
41     inopt = 0
42     maxl = 10
43     maxlrst = 2
44     msbpre = 5
45
46     C * * * * *
47
48     call fnvinit(neq, ier)
49     if (ier .ne. 0) then
50         write(6,1220) ier
51 1220     format('SUNDIALS_ERROR: FNVINIT returned IER = ', i2)
52         stop
```

```

53      endif
54
55      do 20 i = 1, neq
56          uu(i) = 2.0d0 * i
57          scale(i) = 1.0d0
58          constr(i) = 0.0d0
59 20  continue
60
61      call fkinmalloc(msbpre, fnormtol, scsteptol,
62                      &                      constr, inopt, iopt, ropt, ier)
63      if (ier .ne. 0) then
64          write(6,1230) ier
65 1230  format('SUNDIALS_ERROR: FKINMALLOC returned IER = ', i2)
66      call fnvfrees
67      stop
68  endif
69
70      call fkinspgmr(maxl, maxlrst, ier)
71      if (ier .ne. 0) then
72          write(6,1235) ier
73 1235  format('SUNDIALS_ERROR: FKINSPGMR returned IER = ', i2)
74      call fkinfree
75      call fnvfrees
76      stop
77  endif
78
79      call fkinspgmrsetpsol(1, ier)
80      call fkinspgmrsetpset(1, ier)
81
82      write(6,1240)
83 1240  format('Example program kindiagsf:// This fkinsol example code',
84              1           ' solves a 128 eqn diagonal algebraic system.'
85              2           ' Its purpose is to demonstrate the use of the Fortran',
86              3           ' interface'// in a serial environment.'//'
87              4           ' globalstrategy = KIN_INEXACT_NEWTON')
88
89      call fkinsol(uu, globalstrat, scale, scale, ier)
90      if (ier .lt. 0) then
91          write(6,1242) ier, iopt(15)
92 1242  format('SUNDIALS_ERROR: FKINSOL returned IER = ', i2, '/',
93              1           '                           Linear Solver returned IER = ', i2)
94      call fkinfree
95      call fnvfrees
96      stop
97  endif
98
99      write(6,1245) ier
100 1245  format('// FKINSOL return code is ', i3)
101
102      write(6,1246)
103 1246  format('// The resultant values of uu are://')
104
105      do 30 i = 1, neq, 4
106          write(6,1256) i, uu(i), uu(i+1), uu(i+2), uu(i+3)

```

```

107    1256      format(i4, 4(1x, f10.6))
108    30      continue
109
110      write(6,1267) iopt(4), iopt(11), iopt(5), iopt(12), iopt(13),
111      1          iopt(14)
112 1267 format(''Final statistics:''
113      1      ' nni = ', i4, ', nli = ', i4, ', nfe = ', i4,
114      2      ', npe = ', i4, ', nps = ', i4, ', ncfl = ', i4)
115
116      call fkinfree
117      call fnvfrees
118
119      stop
120      end
121
122 c * * * * * * * * * * * * * * * * * * * * * * * * * * * * *
123 c      The function defining the system f(u) = 0 must be defined by a Fortran
124 c      function of the following form.
125
126      subroutine fkfun(uu, fval)
127
128      implicit none
129
130      integer*4 neq, i
131      double precision fval(*), uu(*)
132
133      common /psize/ neq
134
135      do 10 i = 1, neq
136          fval(i) = uu(i) * uu(i) - i * i
137 10      continue
138      return
139      end
140
141
142 c * * * * * * * * * * * * * * * * * * * * * * * * * * * * *
143 c      The routine kpreco is the preconditioner setup routine. It must have
144 c      that specific name be used in order that the c code can find and link
145 c      to it. The argument list must also be as illustrated below:
146
147      subroutine fkpset(udata, uscale, fdata, fscale,
148      1                  vtemp1, vtemp2, ier)
149
150      implicit none
151
152      integer ier
153      integer*4 neq, i
154      double precision pp
155      double precision udata(*), uscale(*), fdata(*), fscale(*)
156      double precision vtemp1(*), vtemp2(*)
157
158      common /pcom/ pp(128)
159      common /psize/ neq
160

```

```

161      do 10 i = 1, neq
162          pp(i) = 0.5d0 / (udata(i) + 5.0d0)
163 10    continue
164      ier = 0
165
166      return
167  end
168
169
170  c * * * * * * * * * * * * * * * * * * * * * * * * * * *
171  c      The routine kpsol is the preconditioner solve routine. It must have
172  c      that specific name be used in order that the c code can find and link
173  c      to it. The argument list must also be as illustrated below:
174
175      subroutine fkpsol(udata, uscale, fdata, fscale,
176                          vv, ftem, ier)
177
178      implicit none
179
180      integer ier
181      integer*4 neq, i
182      double precision pp
183      double precision udata(*), uscale(*), fdata(*), fscale(*)
184      double precision vv(*), ftem(*)
185
186      common /pcom/ pp(128)
187      common /psize/ neq
188
189      do 10 i = 1, neq
190          vv(i) = vv(i) * pp(i)
191 10    continue
192      ier = 0
193
194      return
195  end

```

## D Listing of kindiagpf.f

```
1      program kindiagpf
2      C -----
3      C $Revision: 1.13 $
4      C $Date: 2004/10/15 00:03:58 $
5      C -----
6      C Programmer(s): Allan G. Taylor, Alan C. Hindmarsh and
7      C Radu Serban @ LLNL
8      C -----
9      C Simple diagonal test with Fortran interface, using
10     C user-supplied preconditioner setup and solve routines (supplied
11     C in Fortran, below).
12     C
13     C This example does a basic test of the solver by solving the
14     C system:
15     C       f(u) = 0   for
16     C             f(u) = u(i)^2 - i^2
17     C
18     C       No scaling is done.
19     C       An approximate diagonal preconditioner is used.
20     C
21     C       Execution command: mpirun -np 4 kindiagpf
22     C -----
23     C
24     include "mpif.h"
25
26     integer ier, size, globalstrat, rank, inopt, mype, npes
27     integer maxl, maxlrst
28     integer*4 localsize
29     parameter(localsize=32)
30     integer*4 neq, nlocal, msbpre, baseadd, i, ii
31     integer*4 iopt(40)
32     double precision pp, fnormtol, scsteptol
33     double precision uu(localsize), scale(localsize)
34     double precision constr(localsize)
35
36     common /pcom/ pp(localsize), mype, npes, baseadd, nlocal
37
38     nlocal = localsize
39     neq = 4 * nlocal
40     globalstrat = 1
41     fnormtol = 1.0d-5
42     scsteptol = 1.0d-4
43     inopt = 0
44     maxl = 10
45     maxlrst = 2
46     msbpre = 5
47
48     C       The user MUST call mpi_init, Fortran binding, for the fkinsol package
49     C       to work. The communicator, MPI_COMM_WORLD, is the only one common
50     C       between the Fortran and C bindings. So in the following, the communicator
51     C       MPI_COMM_WORLD is used in calls to mpi_comm_size and mpi_comm_rank
52     C       to determine the total number of processors and the rank (0 ... size-1)
```

```

53   C      number of this process.
54
55   call mpi_init(ier)
56   if (ier .ne. 0) then
57     write(6,1210) ier
58 1210   format('MPI_ERROR: MPI_INIT returned IER = ', i2)
59     stop
60   endif
61
62   call fnvinitp(nlocal, neq, ier)
63   if (ier .ne. 0) then
64     write(6,1220) ier
65 1220   format('SUNDIALS_ERROR: FNVINITP returned IER = ', i2)
66   call mpi_finalize(ier)
67   stop
68   endif
69
70   call mpi_comm_size(MPI_COMM_WORLD, size, ier)
71   if (ier .ne. 0) then
72     write(6,1222) ier
73 1222   format('MPI_ERROR: MPI_COMM_SIZE returned IER = ', i2)
74   call mpi_abort(MPI_COMM_WORLD, 1, ier)
75   stop
76   endif
77
78   if (size .ne. 4) then
79     write(6,1230)
80 1230   format('MPI_ERROR: must use 4 processes')
81   call mpi_finalize(ier)
82   stop
83   endif
84   npes = size
85
86   call mpi_comm_rank(MPI_COMM_WORLD, rank, ier)
87   if (ier .ne. 0) then
88     write(6,1224) ier
89 1224   format('MPI_ERROR: MPI_COMM_RANK returned IER = ', i2)
90   call mpi_abort(MPI_COMM_WORLD, 1, ier)
91   stop
92   endif
93
94   mype = rank
95   baseadd = mype * nlocal
96
97   do 20 ii = 1, nlocal
98     i = ii + baseadd
99     uu(ii) = 2.0d0 * i
100    scale(ii) = 1.0d0
101    constr(ii) = 0.0d0
102 20   continue
103
104   call fkinmalloc(msbpre, fnormtol, scsteptol,
105   &           constr, inopt, iopt, ropt, ier)
106

```

```

107      if (ier .ne. 0) then
108          write(6,1231)ier
109 1231      format('SUNDIALS_ERROR: FKinMalloc returned IER = ', i2)
110      call mpi_abort(MPI_COMM_WORLD, 1, ier)
111      stop
112  endif
113
114      call fkinspgmr(maxl, maxlrst, ier)
115      call fkinspgmrsetpsol(1, ier)
116      call fkinspgmrsetpset(1, ier)
117
118      if (mype .eq. 0) write(6,1240)
119 1240      format('Example program kindiagpf:// This fkinsol example code',
120             1           ' solves a 128 eqn diagonal algebraic system.'
121             2           ' Its purpose is to demonstrate the use of the Fortran',
122             3           ' interface// in a parallel environment.'//
123             4           ' globalstrategy = KIN_INEXACT_NEWTON')
124
125      call fkinsol(uu, globalstrat, scale, scale, ier)
126      if (ier .lt. 0) then
127          write(6,1242) ier, iopt(15)
128 1242      format('SUNDIALS_ERROR: FKINSOL returned IER = ', i2, '/',
129             1           ' Linear Solver returned IER = ', i2)
130      call mpi_abort(MPI_COMM_WORLD, 1, ier)
131      stop
132  endif
133
134      if (mype .eq. 0) write(6,1245) ier
135 1245      format(/' FKINSOL return code is ', i4/)
136
137      if (mype .eq. 0) write(6,1246)
138 1246      format(/' The resultant values of uu (process 0) are://)
139
140      do 30 i = 1, nlocal, 4
141          if(mype .eq. 0) write(6,1256) i + baseadd, uu(i), uu(i+1),
142             1                           uu(i+2), uu(i+3)
143 1256      format(i4, 4(1x, f10.6))
144      30  continue
145
146      if (mype .eq. 0) write(6,1267) iopt(4), iopt(11), iopt(5),
147             1                           iopt(12), iopt(13), iopt(14)
148 1267      format(//Final statistics://
149             1           ' nni = ', i4, ', nli = ', i4, ', nfe = ', i4,
150             2           ' , npe = ', i4, ', nps= ', i4, ', ncfl= ', i4)
151
152      call fkinfree
153      call fnvfreep
154
155  c      An explicit call to mpi_finalize (Fortran binding) is required by
156  c      the constructs used in fkinsol.
157      call mpi_finalize(ier)
158
159      stop
160  end

```

```

161
162
163 C * * * * * * * * * * * * * * * * * * * * * *
164 C      The function defining the system f(u) = 0 must be defined by a Fortran
165 C      function with the following name and form.
166
167      subroutine fkfun(uu, fval)
168
169      implicit none
170
171      integer mype, npes
172      integer*4 baseadd, nlocal, i, localsize
173      parameter(localsize=32)
174      double precision pp
175      double precision fval(*), uu(*)
176
177      common /pcom/ pp(localsize), mype, npes, baseadd, nlocal
178
179      do 10 i = 1, nlocal
180      10     fval(i) = uu(i) * uu(i) - (i + baseadd) * (i + baseadd)
181
182      return
183      end
184
185
186 C * * * * * * * * * * * * * * * * * * * * * *
187 C      The routine kpreco is the preconditioner setup routine. It must have
188 C      that specific name be used in order that the c code can find and link
189 C      to it. The argument list must also be as illustrated below:
190
191      subroutine fkpset(udata, uscale, fdata, fscale,
192                         vtemp1, vtemp2, ier)
193
194      implicit none
195
196      integer ier, mype, npes
197      integer*4 localsize
198      parameter(localsize=32)
199      integer*4 baseadd, nlocal, i
200      double precision pp
201      double precision udata(*), uscale(*), fdata(*), fscale(*)
202      double precision vtemp1(*), vtemp2(*)
203
204      common /pcom/ pp(localsize), mype, npes, baseadd, nlocal
205
206      do 10 i = 1, nlocal
207      10     pp(i) = 0.5d0 / (udata(i)+ 5.0d0)
208
209      ier = 0
210
211      return
212      end
213
214

```

```

215  C * * * * * * * * * * * * * * * * * *
216  C      The routine kpsol is the preconditioner solve routine. It must have
217  C      that specific name be used in order that the c code can find and link
218  C      to it. The argument list must also be as illustrated below:
219
220      subroutine fkpsol(udata, uscale, fdata, fscale,
221                         1                  vv, ftem, ier)
222
223      implicit none
224
225      integer ier, mype, npes
226      integer*4 baseadd, nlocal, i
227      integer*4 localsize
228      parameter(localsize=32)
229      double precision udata(*), uscale(*), fdata(*), fscale(*)
230      double precision vv(*), ftem(*)
231      double precision pp
232
233      common /pcom/ pp(localsize), mype, npes, baseadd, nlocal
234
235      do 10 i = 1, nlocal
236      10     vv(i) = vv(i) * pp(i)
237
238      ier = 0
239
240      return
241      end

```